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(54) Title: MATRIX FOR TRANSDERMAL DRUG DELIVERY

(57) Abstract

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A transdermal drug delivery device involving a macromonomer-containing acrylate or methacrylate copolymer, a softener, and a drug. Also a pressure sensitive skin adhesive involving a macromonomer containing acrylate or methacrylate copolymer and a softener.

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MATRIX FOR TRANSDERMAL DRUG DELIVERY

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Background of the Invention

Field of the Invention

This invention relates to drug containing matrices for use in transdermal drug delivery devices. In another aspect this invention relates to pressure sensitive skin adhesives. In yet another aspect this invention relates to pharmaceutical formulations involving a pressure sensitive skin adhesive layer.

Description of the Related Art

effective amount of drug across the skin of a patient. Devices known to the art include reservoir type devices involving membranes that control the rate of drug release to the skin and devices involving a dispersion of the drug in a matrix.

Certain acrylic copolymers have been used as matrices for delivery of specific drugs. It is critical in such devices that intimate skin contact be achieved and maintained between the skin and the drug-containing matrix. Thus the range of copolymers that are suitable for use as matrices is limited by the ability of the copolymer to comply to the surface of the skin and still release cleanly from the skin. Moreover, the skin presents a substantial barrier to ingress of foreign substances such as drugs into the body. It is therefore often desirable or necessary to incorporate certain materials that enhance the rate at which the drug passes through the skin.

Certain transdermal drug delivery devices have incorporated pressure sensitive adhesive ("PSA") matrices. Fundamentally, PSA's require a balance of viscous and elastic properties which result in a four-fold balance of adhesion, cohesion, stretchiness, and elasticity. In essence, PSA products have sufficient

cohesiveness and elasticity so that, despite their tackiness, they can be handled with the fingers and removed from the skin without leaving substantial residue.

Summary of the Invention

- 5 This invention provides a transdermal drug delivery device, comprising:
 - (1) a backing;

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- (2) a matrix adhered to one side of the backing and comprising
 - (a) a copolymer comprising
- (i) one or more A monomers selected from the group

 consisting of alkyl acrylates containing 4 to 10 carbon atoms in the alkyl group and
 alkyl methacrylates containing 4 to 10 carbon atoms in the alkyl group; and
 - (ii) optionally one or more ethylenically unsaturated B monomers copolymerizable with the A monomer; and
- (iii) a macromonomer, preferably a substantially linear

 macromonomer, copolymerizable with the A and B monomers defined above and having a molecular weight in the range 500-500,000;
 - (b) a softener dissolved in the copolymer; and,
 - (c) if the softener is not therapeutically effective, a therapeutically effective amount of a drug,
 - wherein the structure and amount of the comonomers in the copolymer, the inherent viscosity of the copolymer, and the amount and structure of the drug and the softener are such as to provide the matrix with a compliance value in the range 2×10^{-6} cm²/dyne to about 4×10^{-3} cm²/dyne.
 - It has been found that the copolymer and the softener as defined above can be selected such that the resulting composition adheres to the skin. Accordingly this invention also provides a pressure sensitive skin adhesive comprising:
 - (1) a copolymer comprising
 - (a) one or more A monomers selected from the group consisting of alkyl acrylates containing 4 to 10 carbon atoms in the alkyl group and alkyl methacrylates containing 4 to 10 carbon atoms in the alkyl group; and

(b) optionally one or more ethylenically unsaturated B monomers copolymerizable with the A monomer; and

- (c) a substantially linear macromonomer copolymerizable with the A and B monomers defined above and having a molecular weight in the range 500-500,000; and
 - (2) a softener dissolved in the copolymer,

wherein the structure and amount of the comonomers in the copolymer, the inherent viscosity of the copolymer, and the amount and structure of the softener are such as to provide the pressure sensitive skin adhesive with a compliance value in the range 2×10^{-6} cm²/dyne to about 4×10^{-3} cm²/dyne.

The invention provides a transdermal drug delivery device that allows dissolution of drug and relatively heavy loading with oily excipients, maintains contact with the skin, and can be removed cleanly from the skin. The pressure sensitive skin adhesives of the invention provide these advantages and in addition adhere to the skin.

Detailed Description of the Invention

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The term "lower alkyl" as used herein means straight chain or branched chain alkyl containing 1 to 4 carbon atoms.

The present invention provides a transdermal drug delivery device having a backing and a matrix adhered to one side thereof. It can be adhered directly to a backing or it can be adhered indirectly to a backing via an intermediate layer.

The matrix contains a copolymer as defined above and a softener. The matrix is preferably a pressure sensitive skin adhesive. In addition, the matrix (whether adhesive or not) can be removed cleanly from the skin.

The copolymer utilized in the practice of the invention should be substantially chemically inert to other components utilized in conjugation therewith (e.g., the drugs and/or softeners discussed in detail below). Also the inherent viscosity of the copolymer is such as to ultimately provide a suitable transdermal matrix, preferably a pressure sensitive skin adhesive. Preferably the copolymer has

an inherent viscosity in the range 0.2 dl/g to about 2 dl/g, more preferably in the range 0.4 dl/g to 1.4 dl/g.

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Suitable copolymers comprise one or more A monomers preferably in an amount about 40 to 95 percent by weight, more preferably about 50 to about 70 percent by weight, based on the total weight of all monomers in the copolymer. The A monomer is selected from the group consisting of alkyl acrylates containing 4 to 10 carbon atoms in the alkyl group and alkyl methacrylates containing 4 to 10 carbon atoms in the alkyl group. Examples of suitable alkyl acrylates and methacrylates are n-butyl, n-pentyl, n-hexyl, isoheptyl, n-nonyl, n-decyl, isohexyl, 2-ethyloctyl, isooctyl and 2-ethylhexyl acrylates and methacrylates. Preferred alkyl acrylates include isooctyl acrylate, 2-ethylhexyl acrylate, butyl acrylate, and cyclohexyl acrylate. The most preferred alkyl acrylate is isooctyl acrylate. Preferred alkyl methacrylates include butyl methacrylate, cyclohexyl methacrylate, isobornyl methacrylate, and methyl methacrylate.

The copolymer further optionally comprises one or more ethylenically unsaturated B monomers copolymerizable with the A monomer. Suitable B monomers include those comprising a functional group selected from the group consisting of carboxylic acid, carboxylic acid ester, hydroxy, sulfonamide, urea, carbamate, carboxamide, amine, oxy, oxo, and cyano. The B monomers are preferably used in a total amount from 0 to about 60 percent by weight, more preferably greater than 25 to about 50 percent by weight, and most preferably greater than 30 to about 50 percent by weight (based on the total weight of all the monomers in the copolymer). Preferred B monomers include but are not limited to acrylic acid, methacrylic acid, maleic acid, a hydroxyalkyl acrylate containing 2 to 4 carbon atoms in the hydroxyalkyl group, a hydroxyalkyl methacrylate containing 2 to 4 carbon atoms in the hydroxyalkyl group, acrylamide, methacrylamide, an alkyl substituted acrylamide containing 1 to 8 carbon atoms in the alkyl group, diacetone acrylamide, a dialkyl acrylamide having 1 or 2 carbon atoms in the alkyl group, Nvinyl-N-methyl acetamide, N-vinyl valerolactam, N-vinyl caprolactam, N-vinyl-2pyrrolidone, glycidyl methacrylate, alkoxyethyl acrylate containing 1 to 4 carbon atoms in the alkoxy group, alkoxyethyl methacrylate containing 1 to 4 carbon atoms

in the alkoxy group, 2-ethoxyethoxyethyl acrylate, furfuryl methacrylate, furfuryl acrylate, tetrahydrofurfuryl acrylate, tetrahydrofurfuryl methacrylate, propylene glycol monoacrylate, polyethylene glycol acrylate, polyethylene glycol methyl ether acrylate, polyethylene glycol methacrylate, polyethylene oxide methyl ether acrylate, di(lower)alkylamino ethyl acrylate, di(lower)alkylamino ethyl methacrylate, di(lower)alkylaminopropyl methacrylamide, acrylonitrile, methacrylonitrile, and vinyl acetate.

Particularly preferred B monomers include hydroxyethyl acrylate, acrylamide, hydroxyethyl methacrylate, glyceryl acrylate, N,N-dimethyl acrylamide, 2-ethoxyethoxyethyl acrylate, 2-ethoxyethyl acrylate, tetrahydrofurfuryl acrylate, vinyl acetate and acrylic acid. Most preferred B monomers include hydroxyethyl acrylate and N,N-dimethyl acrylamide, and a combination thereof.

As noted in detail below, the compositions of the invention can contain a relatively high loading of softener. In order to accommodate such loadings the copolymer incorporates a macromonomer, preferably a substantially linear macromonomer, copolymerizable with the A and B monomers defined above and having a molecular weight in the range 500-500,000, preferably 2,000-100,000, and more preferably 5,000-30,000, in an amount (e.g., at least about 0.1 percent by weight based on the total weight of comonomers in the copolymer) effective to control the rheological properties of the copolymer. The macromonomer is generally present in an amount of not more than about 30% by weight based on the total weight of all monomers in the copolymer, more preferably not more than 15%, and most preferably not more than 5%.

The macromonomer can be a compound of the formula

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wherein X is a moiety comprising an ethylenically unsaturated group (such as

-CH₂-C=CH₂, -CH=C(CH₃)(CO₂CH₃), vinyl, or 2-propenyl) copolymerizable with
$$\label{eq:copoly} \begin{picture}(100,0) \put(0.00,0) \put(0$$

the A and B monomers, R² is a hydrogen atom or a lower alkyl group, R³ is a lower alkyl group or the residue of a free-radical initiator, n is an integer from 20 to 500 and each R⁴ is a monovalent radical independently selected from the group consisting of

-CN, and -CO₂R⁶ wherein R⁵ is a hydrogen atom or a lower alkyl group, and R⁶ is a lower alkyl group. Suitable macromonomers include polymethylmethacrylate, styrene/acrylonitrile, and polystyrene macromonomers. Polymethylmethacrylate macromonomers are preferred.

Exemplary macromonomers include those having a general formula selected

20 from the group consisting of

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$$H R^2$$
 $| | |$
 CH_2 -O-C-CH₂(C-CH₂)_nR³
 $| | |$
 $R^7 R^4$

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$$R^{2}$$
 $CH_{2}=C-CH_{2}-(C-CH_{2})_{n}R^{3}$
 $CO_{2}CH_{3}$ R^{4}

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wherein R^7 is a hydrogen atom or a lower alkyl group, R^8 is hydrogen or methyl, and R^2 , R^3 , and R^4 are as defined above.

The macromonomers shown in the formulae directly above are functionally terminated polymers having a single functional group and are sometimes identified as a "semitelechelic" polymers. (Vol. 27 "Functionally Terminal Polymers via Anionic Methods" D. N. Schultz et al., pages 427-440, Anionic Polymerization, American Chemical Society (1981)). Such macromonomers are known and may be prepared by the method disclosed in U.S. Pat. Nos. 3,786,116, 3,842,059 (both to Milkovich et al.), and 4,732,808 (Krampe et al.), the disclosures of which are incorporated herein by reference for the description of the preparation of the macromonomers. Certain macromonomers are commercially available, for example those polymethylmethacrylate macromonomers sold under the trade designation "ELVACITE" by ICI Acrylics (e.g., ELVACITE 1010, a polymethylmethacrylate macromonomer having an inherent viscosity of 0.070-0.080, a T_g of 105°C, a GPC weight average molecular weight of 7,000-10,000, a GPC number average molecular weight of 2,500-4,000, and a polydispersity of 2.5-3.0, and ELVACITE 1020, a polymethylmethacrylate macromonomer having an inherent viscosity of 0.085-0.10, a T₈ of 105°C, a GPC weight average molecular weight of 12,000-15,000, a GPC number average molecular weight of 4,600-6,000, and a polydispersity of 2.5-3.0).

A matrix of the invention further comprises a softener. The softener is dissolved in the matrix. As used herein the term "softener" refers to a generally oily material that raises the compliance value or lowers the glass transition temperature (T_8) of the matrix as compared to the copolymer.

Suitable softeners include certain materials that have been used as skin penetration enhancers or solubilizers in transdermal drug delivery systems.

Exemplary materials include C₈-C₂₂ fatty acids such as isostearic acid, octanoic acid, and oleic acid, C₈-C₂₂ fatty alcohols such as oleyl alcohol and lauryl alcohol, lower alkyl esters of C₈-C₂₂ fatty acids such as ethyl oleate, isopropyl myristate, butyl stearate, and methyl laurate, di(lower) alkyl esters of C₆-C₈ diacids such as diisopropyl adipate, monoglycerides of C₈-C₂₂ fatty acids such as glyceryl monolaurate, tetrahydrofurfuryl alcohol polyethylene glycol ether, polyethylene glycol, propylene glycol, 2-(2-ethoxyethoxy)ethanol, diethylene glycol monomethyl ether, N,N-dimethyldodecylamine-N-oxide, and combinations of the foregoing. Alkylaryl ethers of polyethylene oxide, polyethylene oxide monomethyl ethers, and polyethylene oxide dimethyl ethers are also suitable, as are solubilizers such as dimethyl sulfoxide, glycerol, ethanol, ethyl acetate, acetoacetic ester, N-methyl pyrrolidone, and isopropyl alcohol. Likewise certain drug substances function as softeners, including nicotine, nitroglycerine, chlorpheniramine, nicotinic acid benzyl ester, orphenadrine, scopolamine, and valproic acid.

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Preferred softeners include glyceryl monolaurate, diethylene glycol monomethyl ether, tetrahydrofurfuryl alcohol polyethylene glycol ether, diisopropyl adipate, propylene glycol, isopropyl myristate, ethyl oleate, methyl laurate, 2-(2-ethoxyethoxy)ethanol, and oleyl alcohol.

Preferably the softener is present in not more than that amount which causes the matrix to leave substantial copolymer residue on the skin when peeled from the skin.

While many of the softeners enumerated above are known to affect skin penetration rate, certain softeners affect aspects of performance other than and in addition to skin penetration rate. For example, they are useful in softening or increasing the compliance value and/or lowering the glass transition temperature of otherwise non-compliant (and therefore non-pressure sensitive adhesive) copolymers, rendering them suitable for use as pressure sensitive skin adhesives. However, the softeners enumerated above are generally oily substances that function as plasticizers when incorporated in a copolymer. Such materials can affect adversely the performance of a transdermal matrix, for example by softening it to the point of cohesive failure (where substantial copolymer residue is left on the

skin upon removal of the device from the skin), or by separating from the continuous phase and forming an oily layer that reduces adhesion of an otherwise adhesive matrix. Also, certain softeners (e.g., glyceryl monolaurate, N,N-dimethyldodecylamine-N-oxide) can crystallize in the copolymer, resulting in unstable properties (e.g., unstable drug delivery rates in a transdermal drug delivery device).

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Possible adverse effects of softeners notwithstanding, with proper selection of softeners, monomers and relative amounts thereof, and inherent viscosity of the copolymer, softeners can be included in amounts of up to about 60% by weight based on the total weight of the matrix without cohesive failure or crystal formation, and often without loss of suitable skin adhesion. Softener amounts in excess of 20% and preferably less than about 45% by weight based on the total weight of the matrix have been found to be preferred in order to obtain optimal flux rates in transdermal devices containing the hormone levonorgestrel, and amounts in excess of 30% and less than 45% are more preferred.

The properties desirable in a transdermal matrix are well known to those skilled in the art. For example, it is necessary that the matrix remain in intimate contact with the skin in order to deliver drug at a stable rate. It is desirable for a matrix to have sufficiently little cold flow such that it is stable to flow upon storage. It is also preferred that it release cleanly from the skin, and that it adhere to the skin. In order to achieve skin contact, clean release, preferred levels of adhesion, and resistance to cold flow the amount and structure of the comonomers in the copolymer, the inherent viscosity of the copolymer, and the amount and structure of the softener are selected such that the matrix has a compliance value (measured according to the test method set forth in detail below) in the range 2 x 10⁻⁶ cm²/dyne to about 4 x 10⁻³ cm²/dyne, preferably in the range 3 x 10⁻⁶ cm²/dyne to about 1 x 10⁻³ cm²/dyne and even more preferably in the range 1 x 10⁻³ cm²/dyne to 5 x 10⁻⁴ cm²/dyne. Compliance values outside the broad range recited above sometimes are obtained from materials that are suitable matrices, and even for some that are suitable for use as pressure sensitive skin adhesives. However, those matrices having substantially lower compliance values will generally be relatively

stiff and have less than optimal skin contact and adhesion to skin. Those having substantially higher compliance values will generally have less than optimal cold flow and might leave substantial residue when removed from the skin. Also, a matrix of the invention that is intended for use as a pressure sensitive skin adhesive preferably has a glass transition temperature of -10°C or lower.

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Particularly suitable compositions can be readily selected for a given set of desired properties considering the effects of comonomers, inherent viscosity, and softeners on the properties of the resulting matrix. Certain of such effects are well known to those skilled in the art, and others are described below:

Strongly hydrogen bonding B monomers have been found to increase the amount of polar or hydrogen bonding substances that can be dissolved in a matrix and to decrease the amount of generally nonpolar substances that can be dissolved. Further, a strongly hydrogen bonding copolymer will be a relatively less compliant material. Therefore if B monomers such as acrylic acid or acrylamide are used a lesser amount of macromonomer will be required in order to lower compliance sufficiently to avoid cohesive failure.

Macromonomers also decrease compliance. Therefore a given target compliance value can often be achieved using a lower inherent viscosity A/B copolymer combination and a greater amount of macromonomer, or a higher inherent viscosity A/B combination and less macromonomer.

A relatively high compliance pressure sensitive skin adhesive involving a macromonomer will generally have better adhesive properties than an A/B copolymer having the same compliance value. Increasing macromonomer content generally increases the amount of softener that can be loaded into a pressure sensitive skin adhesive without cohesive failure. Increasing inherent viscosity will also tend to allow higher softener loading without cohesive failure.

A change that would increase inherent viscosity of a copolymer (such as increased molecular weight through selection of polymerization conditions and/or solvent ratios) will generally decrease compliance.

Further conventional components, such as stabilizers and reinforcers (e.g., colloidal silicon dioxide), can be incorporated into the matrix if necessary or desirable.

Of course such high levels of certain individual softeners (e.g., N,N
dimethyldodecylamine-N-oxide) are to be avoided in order to avoid excessive skin irritation.

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The matrix of a transdermal drug delivery device of the invention further comprises a drug. Suitable drugs include those active substances enumerated above in connection with softeners, as well as antiinflammatory drugs, both steroidal (e.g., hydrocortisone, prednisolone, triamcinolone) and nonsteroidal (e.g., naproxen, piroxicam); antibacterials (e.g., penicillins such as penicillin V, cephalosporins such as cephalexin, erythromycin, tetracycline, gentamycin, sulfathiazole, nitrofurantoin, and quinolones such as norfloxacin, flumequine, and ibafloxacin); antiprotazoals (e.g., metronidazole); antifungals (e.g., nystatin); coronary vasodilators (e.g., nitroglycerin); calcium channel blockers (e.g., nifedipine, diltiazem); bronchodilators (e.g., theophylline, pirbuterol, salmeterol, isoproterenol); enzyme inhibitors such as collagenase inhibitors, protease inhibitors, elastase inhibitors, lipoxygenase inhibitors (e.g., A64077), and angiotensin converting enzyme inhibitors (e.g., captopril, lisinopril); other antihypertensives (e.g., propranolol); leukotriene antagonists (e.g., ICI204,219); anti-ulceratives such as H2 antagonists; steroidal hormones (e.g., progesterone, testosterone, estradiol, levonorgestrel); antivirals and/or immunomodulators (e.g., 1-isobutyl-1H-imidazo[4,5-c]quinolin-4-amine, 1-(2-hydroxy-2-methylpropyl)-1H-imidazo[4,5-c]quinoline-4-amine, and other compounds disclosed in U.S. Pat. No. 4,689,338, incorporated herein by reference, acyclovir); local anesthetics (e.g., benzocaine, propofol); cardiotonics (e.g., digitalis, digoxin); antitussives (e.g., codeine, dextromethorphan); antihistamines (e.g., diphenhydramine, chlorpheniramine, terfenadine); narcotic analgesics (e.g., morphine, fentanyl); peptide hormones (e.g., human or animal growth hormones, LHRH); cardioactive products such as atriopeptides; proteinaceous products (e.g., insulin); enzymes (e.g., anti-plaque enzymes, lysozyme, dextranase); antinauseants (e.g., scopolomine); anticonvulsants (e.g., carbamazine); immunosuppressives (e.g.,

cyclosporine); psychotherapeutics (e.g., diazepam); sedatives (e.g., phenobarbital); anticoagulants (e.g., heparin); analgesics (e.g., acetaminophen); antimigraine agents (e.g., ergotamine, melatonin, sumatriptan); antiarrhythmic agents (e.g., flecainide); antiemetics (e.g., metaclopromide, ondansetron); anticancer agents (e.g., methotrexate); neurologic agents such as anxiolytic drugs; hemostatics; anti-obesity agents; and the like, as well as pharmaceutically acceptable salts and esters thereof.

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The drug is present in a transdermal delivery device of the invention in a therapeutically effective amount, i.e., an amount effective to bring about a desired therapeutic result in the treatment of a condition. The amount that constitutes a therapeutically effective amount varies according to the particular drug incorporated in the device, the condition being treated, any drugs being coadministered with the selected drug, desired duration of treatment, the surface area of the skin over which the device is to be placed, and other components of the transdermal delivery device. Accordingly it is not practical to enumerate particular preferred amounts but such can be readily determined by those skilled in the art with due consideration of these factors. Generally, however, a drug is present in a transdermal device of the invention in an amount of about 0.01 to about 30 percent by weight based on the total weight of the matrix. In a preferred embodiment the drug is substantially fully dissolved, and the matrix is substantially free of solid undissolved drug.

A transdermal delivery device or an adhesive coated sheet material of the invention also comprises a backing. The backing is flexible such that the device conforms to the skin. Suitable backing materials include conventional flexible backing materials used for pressure sensitive tapes, such as polyethylene, particularly low density polyethylene, linear low density polyethylene, high density polyethylene, polyester, polyethylene terephthalate, randomly oriented nylon fibers, polypropylene, ethylene-vinyl acetate copolymer, polyurethane, rayon and the like. Backings that are layered, such as polyethylene-aluminum-polyethylene composites, are also suitable. The backing should be substantially inert to the ingredients of the matrix layer.

The copolymers described above for use in a device of the invention can be prepared by methods well known to those skilled in the art and described, for example, in U.S. Patent RE 24,906 (Ulrich) and U.S. Pat. No. 4,732,808 (Krampe at al.), the disclosures of which are incorporated herein by reference.

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Matrices of the invention can be used in the form of an adhesive coated sheet material. Such sheet materials are preferably prepared by combining the copolymer, the softener, and any additional components (e.g., a drug) with an organic solvent (e.g., ethyl acetate, methanol, acetone, 2-butanone, ethanol, isopropyl alcohol, toluene, alkanes, or a mixture thereof) to afford a coating formulation. The total solids content of the coating formulation is preferably in a range of about 15 to 40 percent by weight, and more preferably in the range of about 20 to 35 percent by weight, based on the total weight of the coating formulation. The components of the coating formulation are combined and mixed (e.g., by shaking or rolling) until a homogeneous formulation is obtained, then allowed to stand to dissipate air bubbles. The resulting coating formulation is knife coated onto a suitable release liner to provide a predetermined uniform thickness of the coating formulation. Suitable release liners include conventional release liners comprising a known sheet material such as a polyester web, a polyethylene web, or a polystyrene web, or a polyethylene-coated paper, coated with a suitable fluoropolymer or silicone based coating. The coated release liner is dried and then laminated onto a backing material using conventional methods. Alternatively the coating formulation can be coated directly onto a backing. A transdermal device involving a matrix that is not a skin adhesive can be fixed to the skin by conventional means such as a peripheral ring of a pressure sensitive skin adhesive.

Adhesive coated sheet materials of the invention can be made in the form of an article such as a tape, a patch, a sheet, a dressing or any other form known to those skilled in the art. Transdermal drug delivery devices generally are made in the form of a patch of a size suitable to deliver a preselected amount of a drug through the skin. Generally the transdermal device will have a surface area of about 1 cm² to about 40 cm².

The examples set forth below are intended to illustrate the invention.

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Compliance Test Method

The compliance values given in the examples below were obtained using a modified version of the Creep Compliance Procedure described in U.S. Pat. No. 4,737,559 (Kellen), the disclosure of which is incorporated herein by reference. The release liner is removed from a sample of the material to be tested. The exposed adhesive surface is folded back on itself in the lengthwise direction to produce a "sandwich" configuration, i.e., backing/adhesive/backing. The "sandwiched" sample is passed through a laminator, or alternatively rolled with a hand-operated roller, then two test samples of equal area are cut using a rectangular die. One test sample is centered on a first stationary plate of a shear-creep rheometer with the long axis of the test sample centered on the short axis of the plate. The small, non-stationary plate of the shear-creep rheometer is centered over the first sample on the first stationary plate such that the hook is facing up and toward the front of the rheometer. The second test sample is centered on the upper surface of the small, non-stationary plate matching the axial orientation of the first test sample. A second stationary plate is placed over the second test sample and the entire assembly is clamped into place. The end of the small, non-stationary plate that is opposite the end with the hook is connected to a chart recorder. A string is connected to the hook of the small, non-stationary plate and extended over the front pulley of the rheometer. A weight (e.g., 500 g) is attached to the free end of the string. The chart recorder is started and at the same time the weight is quickly released so that it hangs free. The weight is removed after exactly 3 minutes has elapsed. The displacement is read from the chart recorder. The compliance is then calculated using the equation:

$$J=2\frac{AX}{hf}$$

where A is the area of one face of the test sample, h is the thickness of the adhesive mass (i.e., two times the matrix thickness of the sample being tested), X is the displacement and f is the force due to the mass attached to the string. Where A is

expressed in cm², h in cm, X in cm and f in dynes, the compliance value is given in cm²/dyne.

Determination of Isopropyl Myristate Content

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The amount of isopropyl myristate present in a pressure sensitive skin adhesive composition was determined using the following test method. The release liner is removed from a sample of the material to be tested. The adhesive coating is manually scraped from the backing film. A 15 mg portion of the adhesive coating is placed into a clean sample vial. Tetrahydrofuran (2 mL containing 0.10 mg/mL of lauryl acrylate which serves as an internal standard) is added and the sample is mixed until all of the adhesive coating is dissolved. A portion of the solution is placed in an autosampler vial and analyzed by gas chromatography using the following conditions: Instrument: HP5890, Column: DB-5, 30 meter, 0.25 µM film, 0.25 mm I.D.; Temperature Program: Initial 100°C, ramp 10°C/min to 300°C, hold 2 min; Injection: 2 µL, split 25/1, 300°C; Detection: FID, 300°C. Isopropyl myristate standards are prepared using copolymer samples containing no isopropyl myristate. Separate standard curves are prepared for each copolymer. Each sample is run in duplicate.

Determination of Oleyl Alcohol Content

The amount of oleyl alcohol present in a pressure sensitive skin adhesive composition was determined using the following test method. The release liner is removed from a sample of the material to be tested. The adhesive coating is manually scraped from the backing film. A 15 mg portion of the adhesive coating is placed into a clean sample vial. Tetrahydrofuran (10 mL containing 0.1 mg/mL of dodecyl alcohol which serves as an internal standard) is added and the sample is mixed until all of the adhesive coating is dissolved. A portion of the solution is placed in an autosampler vial and analyzed by gas chromatography using the following conditions: Instrument: HP5890; Column: DB-wax, 15 meter, 0.25 μM film, 0.25 mm I.D.; Temperature Program: Initial 60°C, ramp 7°C/min to 250°C, hold 2 min; Injection: 2 μL, split 25/1, 250°C; Detection: FID, 250°C. Oleyl

alcohol standards are prepared using copolymer samples containing no oleyl alcohol. Separate standard curves are prepared for each copolymer. Each sample is run in duplicate.

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Preparation of Copolymers

The copolymers used in the examples that follow were prepared generally according to the methods described below. The inherent viscosity values which are reported were measured by conventional means using a Cannon-Fenske #50 viscometer in a water bath controlled at 27°C to measure the flow time of 10 milliliters of a polymer solution (0.15-0.25 g per deciliter of polymer in ethyl acetate, unless other wise indicated). The test procedure followed and the apparatus used are described in detail in "Textbook of Polymer Science", F. W. Billmeyer, Wiley Interscience, Second Edition, 1971, Pages 84 and 85.

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Preparation of Isooctyl Acrylate: Dimethylacrylamide:
Hydroxyethyl Acrylate: Polymethylmethacrylate Macromonomer
(60/15/15/10) Copolymer

Isooctyl acrylate (141.0 g), N,N-dimethylacrylamide (35.25 g), hydroxyethyl acrylate (35.25 g), ELVACITE^{IM} 1010 polymethylmethacrylate macromonomer (23.50 g, ICI), ethyl acetate (251.75 g), isopropanol (13.25 g) and 2,2'-azobis(2,4-dimethylpentanenitrile) (0.47 g, VAZO^{IM} 52 available from DuPont) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1L/min) for 2 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed, opened, charged with an additional 0.47 g of VAZO 52, repurged with nitrogen as before, sealed and placed in the launderometer for an additional 24 hours. The percent solids of the resulting solution of copolymer was 45.51%. The inherent viscosity was 0.469 deciliter/gram in ethyl acetate at 0.25 g/dl.

Preparation of Isooctyl Acrylate:

Dimethylacrylamide: Polymethylmethacrylate Macromonomer (50/40/10) Copolymer

Isooctyl acrylate (117.5 g), N,N-dimethylacrylamide (94.0 g), ELVACITE^M

1010 polymethylmethacrylate macromonomer (23.5 g), ethyl acetate (251.75 g), isopropanol (13.25 g) and VAZO 52 (0.47 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1L/min) for 2 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed, opened, charged with an additional 0.47 g of VAZO 52, repurged with nitrogen as before, sealed and placed in the launderometer for an additional 24 hours. The percent solids of the resulting solution of copolymer was 46.19%. The inherent viscosity was 0.532 dl/g in ethyl acetate at 0.25 g/dl.

Preparation of Isooctyl Acrylate:

Dimethylacrylamide: Polymethylmethacrylate Macromonomer (63/27/10) Copolymer

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Isooctyl acrylate (157.5 g), N,N-dimethylacrylamide (67.5 g), ELVACITE 1010 macromonomer (25.0 g), ethyl acetate (261.25 g), isopropanol (13.75 g) and VAZO 52 (0.5 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed, opened, charged with an additional 0.5 g of VAZO 52, repurged with nitrogen as before, sealed and placed in the launderometer for an additional 24 hours. The percent solids of the resulting solution of copolymer was 47.8%. The inherent viscosity was 0.394 dl/g in ethyl acetate at 0.15 g/dl.

Preparation of Isooctyl Acrylate: Hydroxyethyl Acrylate: Polymethylmethacrylate

Macromonomer

(55/40/5) Copolymer

Molecular sieves (50 g of 8-12 mesh, 4A, 1.6 mm beads) were added to each of 4 quart (0.95 L) wide mouth jars. The jars were filled with isooctyl acrylate, hydroxyethyl acrylate, ethyl acetate, and isopropanol respectively. The jars were tightly capped and allowed to stand for at least 24 hours. The molecular sieves were then removed by filtration through Whatman filter paper No. 4. The "dry" monomers and solvents were then stored in tightly capped bottles until used to prepare copolymer. Isooctyl acrylate (137.5 g), hydroxyethyl acrylate (100.0 g), ELVACITETM 1010 polymethylmethacrylate macromonomer (12.5 g), ethyl acetate (318.75 g), isopropanol (56.25 g) and VAZO 52 (0.5 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed, opened, charged with an additional 0.5 g of VAZO 52, repurged with nitrogen as before, sealed and placed in the launderometer for an additional 24 hours. The percent solids of the resulting solution of copolymer was 39.30%. The inherent viscosity was 0.335 dl/g in ethyl acetate at 0.15 g/dl.

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Preparation of Isooctyl Acrylate:

Hydroxyethyl acrylate: Polystyrene Macromonomer (54/36/10) Copolymer

Isooctyl acrylate (135 g), hydroxyethyl acrylate (90 g), polystyrene 25 macromonomer (25.0 g), ethyl acetate (356.25 g), isopropanol (18.75 g) and VAZO 52 (0.5 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed, opened, charged with an additional 0.5 g of VAZO 52, repurged with nitrogen as before, sealed and placed in the launderometer for an additional 24

hours. The percent solids of the resulting solution of copolymer was 41.2%. The inherent viscosity was 0.75 dl/g in ethyl acetate at 0.15 g/dl.

Preparation of Isooctyl Acrylate:

Hydroxyethyl acrylate: Polystyrene Macromonomer (54/36/10) Copolymer

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Isooctyl acrylate (135 g), hydroxyethyl acrylate (90 g), polystyrene macromonomer (25.0 g), ethyl acetate (318.75 g), isopropanol (56.25 g) and VAZO 52 (0.5 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed, opened, charged with an additional 0.5 g of VAZO 52, repurged with nitrogen as before, sealed and placed in the launderometer for an additional 24 hours. The percent solids of the resulting solution of copolymer was 39.6%. The inherent viscosity was 0.29 dl/g in ethyl acetate at 0.15 g/dl.

Preparation of Isooctyl Acrylate:Polystyrene Macromonomer (95/5) Copolymer

Isooctyl acrylate (237.5 g), polystyrene macromonomer (12.5 g), ethyl acetate (261.25 g), isopropanol (13.75 g) and VAZO 52 (0.5 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed, opened, charged with an additional 0.5 g of VAZO 52, repurged with nitrogen as before, sealed and placed in the launderometer for an additional 24 hours. The percent solids of the resulting solution of copolymer was 47.5%. The inherent viscosity was 0.45 dl/g in ethyl acetate at 0.15 g/dl.

Preparation of Isooctyl Acrylate:

Vinyl Acetate: Polystyrene Macromonomer

(61/37/2) Copolymer

Isooctyl acrylate (134.2 g), vinyl acetate (81.4 g), polystyrene

5 macromonomer (4.4 g), 2,2'-azobis(isobutyronitrile) (0.55 g), ethyl acetate (126.0 g), and toluene (54.0 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1 L/min) for 2 minutes. The bottle was sealed and placed in a rotating water bath at 60°C for 24 hours. The resulting copolymer solution was diluted with ethyl acetate (150 mL). The inherent viscosity in ethyl acetate at 0.2 g/dl was measured at 0.87 dl/g.

Preparation of Isooctyl Acrylate:

Vinyl Acetate: Polystyrene Macromonomer (61/37/2) Copolymer

Isooctyl acrylate (134.2 g), vinyl acetate (81.4 g), polystyrene macromonomer (4.4 g), 2,2'-azobis(isobutyronitrile) (0.55 g), ethyl acetate (144.0 g), and toluene (36.0 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1 L/min) for 2 minutes. The bottle was sealed and placed in a rotating water bath at 60°C for 24 hours. The resulting copolymer solution was diluted with ethyl acetate (150 mL). The inherent viscosity in ethyl acetate at 0.2 g/dl was measured at 1.02 dl/g.

Preparation of Isooctyl Acrylate:

Vinyl Acetate: Polystyrene Macromonomer

(58/37/5) Copolymer

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Isooctyl acrylate (127.6 g), vinyl acetate (81.4 g), polystyrene macromonomer (11.0 g), 2,2'-azobis(isobutyronitrile) (0.55 g), ethyl acetate (126.0), and toluene (54.0 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1 L/min) for 2 minutes. The bottle was sealed and placed in a rotating water bath at 60°C for 24 hours. The resulting

copolymer solution was diluted with ethyl acetate (150 mL). The inherent viscosity in ethyl acetate at 0.2 g/dl was measured at 0.89 dl/g.

Preparation of Isooctyl Acrylate:

Vinyl Acetate: Polystyrene Macromonomer (58/37/5) Copolymer

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Isooctyl acrylate (127.6 g), vinyl acetate (81.4 g), polystyrene macromonomer (11.0 g), 2,2'-azobis(isobutyronitrile) (0.55 g), ethyl acetate (144.0), and toluene (36.0 g) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1 L/min) for 2 minutes. The bottle was sealed and placed in a rotating water bath at 60°C for 24 hours. The resulting copolymer solution was diluted with ethyl acetate (150 mL). The inherent viscosity in ethyl acetate at 0.2 g/dl was measured at 1.02 dl/g.

Preparation of Isooctyl Acrylate:

Vinyl Acetate: Polymethylmethacrylate Macromonomer (58/37/5) Copolymer

Isooctyl acrylate (145.0 g), vinyl acetate (92.5 g), ELVACITE™ 1020 polymethylmethacrylate macromonomer (12.5 g), 2,2'-azobis(2,4-dimethylpentanenitrile) (0.5 g), and ethyl acetate (282.0) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1 L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed from the bath, opened, charged with an additional 0.5 g of 2,2'-azobis(2,4-dimethylpentanenitrile), deoxygenated as before, sealed and returned to the rotating water bath for an additional 24 hours. The inherent viscosity in ethyl acetate at 0.15 g/dl was measured at 1.05 dl/g.

Preparation of Isooctyl Acrylate:

Vinyl Acetate: Polymethylmethacrylate Macromonomer (58/37/5) Copolymer

Isooctyl acrylate (145.0 g), vinyl acetate (92.5 g), ELVACITE™ 1020

5 polymethylmethacrylate macromonomer (12.5 g), 2,2'-azobis(2,4-dimethylpentanenitrile) (0.5 g), and ethyl acetate (250.0) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1 L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed from the bath, opened, charged with an additional 0.5 g of 2,2'-azobis(2,4-dimethylpentanenitrile), deoxygenated as before, sealed and returned to the rotating water bath for an additional 24 hours. The inherent viscosity in ethyl acetate at 0.15 g/dl was measured at 1.15 dl/g.

Preparation of Isooctyl Acrylate:

Vinyl Acetate: Polymethylmethacrylate Macromonomer (53/37/10) Copolymer

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Isooctyl acrylate (132.5 g), vinyl acetate (92.5 g), ELVACITE™ 1020 polymethylmethacrylate macromonomer (25.0 g), 2,2'-azobis(2,4-dimethylpentanenitrile) (0.5 g), and ethyl acetate (230.8) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1 L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed from the bath, opened, charged with an additional 0.5 g of 2,2'-azobis(2,4-dimethylpentanenitrile), deoxygenated as before, sealed and returned to the rotating water bath for an additional 24 hours. The inherent viscosity in ethyl acetate at 0.15 g/dl was measured at 0.815 dl/g.

Preparation of Isooctyl Acrylate:

Vinyl Acetate: Polymethylmethacrylate Macromonomer (53/37/10) Copolymer

Isooctyl acrylate (132.5 g), vinyl acetate (92.5 g), ELVACITETM 1020 polymethylmethacrylate macromonomer (25.0 g), 2,2'-azobis(2,4-

dimethylpentanenitrile) (0.5 g), and ethyl acetate (204.5) were charged into a one liter bottle. The mixture was deoxygenated by purging with nitrogen (1 L/min) for 3 minutes. The bottle was sealed and placed in a rotating water bath at 45°C for 24 hours. The bottle was removed from the bath, opened, charged with an additional 0.5 g of 2,2'-azobis(2,4-dimethylpentanenitrile), deoxygenated as before, sealed and returned to the rotating water bath for an additional 24 hours. The inherent viscosity in ethyl acetate at 0.15 g/dl was measured at 0.92 dl/g.

Preparation of "Dried" Adhesive

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Dried adhesive is prepared by knife coating a 25 to 50 percent solids solution of the adhesive copolymer at a thickness of 20 to 25 mil (500 to 635 μ M) onto a release liner. The adhesive coated release liner is oven dried (e.g. 4 min at 110°F (43°C), 2 minutes at 185°F (85°C), and 10 minutes at 300°F (149°C)) to remove solvent and reduce the amount of residual monomers. The dried adhesive copolymer is stripped off the release liner and stored in a glass container.

In the examples that follow all percentages are weight/weight unless otherwise indicated. The weight percentages of the formulations after drying are calculated values, unless otherwise indicated, and assume that only solvent was evaporated during the drying process. The abbreviations IOA, HEA, DMACM, PSMac, PMMAMac, and VoAc are used for isooctyl acrylate, hydroxyethyl acrylate, dimethylacrylamide, polystyrene macromonomer, polymethylmethacrylate macromonomer, and vinyl acetate respectively. The polystyrene macromonomer used in the copolymers in the examples below is that macromonomer designated as Example M-1 in U.S. Pat. No. 4,732,808 (Krampe). Except as noted, the polymethylmethacrylate macromonomer used was ELVACITE 1010. The abbreviations BS, DDAO, DGME, DIPA, EO, GML, IPM, ISA, LG, ML, OA and PG are used for butyl stearate, N,N-dimethyldodecylamine-N-oxide, diethylene glycol monoethyl ether, diisopropyl adipate, ethyl oleate, glyceryl monolaurate, isopropyl myristate, isostearic acid, lauryl glycol, methyl laurate, oleyl alcohol and propylene glycol respectively. The abbreviation LN is used for levonorgestrel.

Example 1

Copolymer (50 g of 54/36/10 IOA/HEA/PSMac, 41% solids in 95/5 ethyl acetate/isopropanol, inherent viscosity ("iv") = 0.75 dl/g) and isopropyl myristate (1.08 g) were combined in a glass jar. The jar was capped and placed on a roller for about 24 hours. The resulting formulation was knife coated at a wet thickness of 12 mil (305 μM) onto a silicone release liner [5 mil (127 μM) Daubert PESTER]. The coated release liner was oven dried at 110°F (43°C) for 4 minutes then at 180°F (82°C) for 4 minutes. The resulting coating contained 95 percent 54/36/10 IOA/HEA/PSMac copolymer and 5 percent isopropyl myristate. The coated liner was laminated to the corona treated side of a 3 mil (76 μM) polyethylene film. The compliance was measured using the test method described above and found to be 0.42 X 10⁻⁵ cm²/dyne (average of three independent determinations).

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Examples 2 - 33

Using the general method of Example 1, a series of coated sheet materials in which the copolymer, softener and amount of softener were varied was prepared. The copolymer, identity and amount of softener, wet coating thickness, and the compliance values are shown in Table 1. Unless otherwise indicated, each J-value is the average of three independent determinations. When the compliance was "not run", the formulation was too soft to be tested.

		Table 1			
Example Number	Copolymer		Softener	Wet Coating Thickness (mil/μΜ)	J-value (X 10° ⁵ cm²/dyne)
	Туре	iv (dVg)			
2	54/36/10 IOA/HEA/PSMac	0.75	10% IPM	12/305	0.57
3	54/36/10 IOA/HEA/PSMac	0.75	13% IPM	12/305	0.57
4	54/36/10 IOA/HEA/PSMac	0.75	17% IPM	10/254	0.80
\$	54/36/10 IOA/HEA/PSMac	0.75	20% IPM	10/254	1.12
9	54/36/10 IOA/HEA/PSMac	0.75	25% IPM	8/203	2.26
7	54/36/10 IOA/HEA/PSMac	0.29	5% IPM	12/305	1.09
∞	54/36/10 IOA/HEA/PSMac	0.29	10% IPM	12/305	1.65
6	54/36/10 IOA/HEA/PSMac	0.29	13% IPM	12/305	1.83
10	54/36/10 IOA/HEA/PSMac	0.29	17% IPM	10/254	2.131

		Table 1			
Example Number	Copolymer		Softener	Wet Coating Thickness (mil/µM)	J-value (X 10 ⁻⁵ cm ² /dvne)
· <u></u>	Type	iv (d//g)			
11	54/36/10 IOA/HEA/PSMac	0.29	20% IPM	10/254	3.87²
12	54/36/10 IOA/HEA/PSMac	0.29	25% IPM	8/203	14.2
13	51/34/15 IOA/HEA/PMMAMac	0.38	10% IPM	12/305	0.28
14	51/34/15 IOA/HEA/PMMAMac	0.38	20% IPM	10/254	0.46
15	51/34/15 IOA/HEA/PMMAMac*	0.42	10% IPM	12/305	0.28
16	51/34/15 IOA/HEA/PMIMAMac*	0.42	20% IPM	10/254	0.38
17	72/13/15 IOA/HEA/PMIMAMac	0.36	10% IPM	12/305	0.38
18	72/13/15 IOA/HEA/PMIMAMac	0.36	20% IPM	10/254	0.53
61	85/15 IOA/PMMAMac	0.48	10% IPM	12/305	not run

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		Table 1			
Example Number	Copolymer		Softener	Wet Coating Thickness (mil/μΜ)	J-value (X 10° ⁵ cm²/dyne)
	Type	iv (dVg)			
20	85/15 IOA/PMIMAMac	0.48	20% IPM	10/254	off scale
CI	57/38/5 IOA/HEA/PSMac	0.32	попе	6/152	1.29
21	54/36/10 IOA/HEA/PSMac	0.29	30% IPM	6/152	8.99
22	51/34/15 IOA/HEA/PSMac	0.28	30% IPM	6/152	18.2
23	\$1/34/15 IOA/HEA/PSMac	0.28	15% IPM	10/254	0.76
CZ	57/38/5 IOA/HEA/PSMac	0.65	none	6/152	0.57
24	54/36/10 IOA/HEA/PSMac	0.75	35% IPM	6/152	11.2
25	\$1/34/15 IOA/HEA/PSMac	0.73	50% IPM	6/152	551
26	51/34/15 IOA/HEA/PSMac	0.73	40% IPM	6/152	27.8

		Table 1			
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Example	Copolymer		Softener	Wet Coating	J-value
				Thickness	(X 10 ⁻⁵
				(mil/µM)	cm²/dyne)
	Туре	٧i		` 	
		(dl/g)			
27	51/34/15 IOA/HEA/PSMac	0.73	30% IPM	6/152	2.36
28	51/34/15 IOA/HEA/PSMac	0.73	50% OA	10/254	100
29	51/34/15 IOA/HEA/PSMac	0.73	400,004		lint los
		2	40% UA	10/254	3.59
30	51/34/15 IOA/HEA/PSMac	0.73	30% OA	10/254	0.64
31	51/34/15 IOA/HEA/PSMac	0.73	20% OA	10/254	0.42
32	51/34/15 IOA/HEA/PSMac	0.73	40% ISA	10/254	0.79
33	51/34/15 IOA/HEA/PSMac	0.73	40% BS	10/254	nin Jon

¹average of 2 determinations
²average of 4 determinations
PMMAMac* ELVACITE 1020

Examples 34 - 38

Using the general method of Example 1, a series of coated sheet materials in which the copolymer was varied but the amount of IPM was theoretically held constant was prepared. The copolymer and amount (both calculated and determined using a modification of the method described above) of IPM, wet coating thickness, and the compliance values are shown in Table 2. In the modified analysis procedure, sample preparation involved combining 2 mL ethyl acetate containing 0.05 mg/mL lauryl acrylate with 25 mg of polymer. In the modified analysis procedure, isopropyl myristate standards did not contain copolymer.

Unless otherwise indicated, each J-value is the average of three independent determinations.

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	J-value (X 10 ⁻⁵	cm²/dyne)			1.68²	3.86	12.8	19.7	10.3
	Wet Coating Thickness	(mil/µM)			10/254	10/254	10/254	10/254	10/254
	Wt Percent IPM		Actual		13.5	11.7	12.5	13.4	10.5
	Wt Perc		Calc.		20	20	20	20	20
Table 2			i,	(dl/g)	1.601	1.071	0.47	0.38	0.34
	Copolymer		Туре		78/14/8 IOA/HEA/PSMac	78/14/8 IOA/HEA/PSMac	95/5 IOA/PSMac	55/40/5 IOA/HEA/PSMac	55/40/5 IOA/HEA/PMIMAMac
	Example Number				34	35	36	37	38

Run in tetrahydrofuran

²Average of 4 determinations

Example 39

Copolymer (50 g of 51/34/15 IOA/HEA/PSMac, 39.2% solids in 95/5 ethyl acetate/isopropanol, iv = 0.73 dl/g) and oleyl alcohol (8.4 g) were combined in a glass jar. The jar was capped and placed on a roller for about 24 hours. The resulting formulation was knife coated at a wet thickness of 15 mil (381 μM) onto a silicone release liner [5 mil (127 μM) Daubert PESTER]. The coated release liner was oven dried at 110°F (43°C) for 20 minutes. The resulting coating theoretically contained 70 percent 51/34/15 IOA/HEA/PSMac copolymer and 30 percent oleyl alcohol. The coated liner was laminated to a backing (1109 SCOTCHPAKTM tan, polyester film laminate, available from the 3M Company). The compliance was measured using the test method described above and found to be 0.74 X 10⁻⁵ cm²/dyne (average of three independent determinations). A portion of the coating was removed from the backing and assayed for oleyl alcohol using the test method described above. The oleyl alcohol content was found to be 28 percent.

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Examples 40 - 106

Using the general method of Example 39, a series of coated sheet materials in which the copolymer, softener and amount of softener were varied was prepared. The copolymer, identity and amount (weight percent, both calculated and determined using the methods described above) of softener, wet coating thickness, and the compliance values are shown in Table 3. Unless otherwise indicated, each J-value is the average of three independent determinations.

		Table 3				
Copolymer	į		Softener	_	Wet Coating Thickness (mil/µM)	J-value (X 10 ⁻⁵ cm²/dyne)
Type	vi (g/lb)	Д	Calc	Actual		
57/38/5 IOA/HEA/PSMac	0.65	None	0	0	18/381	0.92
57/38/5 IOA/HEA/PSMac	0.65	0A	10	8.9	15/381	2.05
57/38/5 IOA/HEA/PSMac	0.65	0A	20	19.9	18/381	3.39
57/38/5 IOA/HEA/PSMac	0.65	0A	30	29.7	15/381	4.29
95/5 IOA/PSMac	0.45	none	0	0	18/381	3.22
95/5 IOA/PSMac	0.45	OA	20	18.9	15/381	\$.00
95/5 IOA/PSMac	0.45	0A	40	37.1	182/31	8.16
90/10 IOA/PSMac	0.65	none	0	0	15/381	1.07

Table 3	Softener Wet Coating J-value Thickness (X 10 ⁻⁵ (mil/µM) cm²/dyne)	iv ID Calc Actual (dVg)	0.65 OA 20 18.8 15/381 1.63	0.65 OA 40 39 15/381 2.72	0.55 none 0 0 15/381 0.56	0.55 OA 20 19 15/381 0.85	0.55 OA 40 36 15/381 1.74	0.65 OA 40 37 15/381 4.99	0.65 OA 60 56.5 15/381 221 ²	0.65 OA 60 - 4/102 1300 ²	
Table 3	Softener	<u> </u>	20	40	0	20	40	40	09	09	9,0
		О	0A	OA	none	V	OA	V 0	Φ0	Φ0	
		vi (dVB)	0.65	9.65	0.55	0.55	0.55	0.65	0.65	0.65	3, 6
	Copolymer	Type	90/10 IOA/PSMac	90/10 IOA/PSMac	85/15 IOA/PSMac	85/15 IOA/PSMac	85/15 IOA/PSMac	57/38/5 IOA/HEA/PSMac	57/38/5 IOA/HEA/PSMac	57/38/5 IOA/HEA/PSMac	2 10 4 1 0 2 2 2 0
	Example Number		45	46	9)	47	48	49	50	51	2

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	J-value (X 10 ⁻⁵ cm ² /dyne)		not run	not run	2.95	not run	4.121	1.99	48.22	2.82³	0.51
	Wet Coating Thickness (mil/µM)		15/381	4/102	15/381	15/381	4/102	15/381	15/381	4/102	15/381
		Actual	52.8	•	38	56.6		40.5	09		0
	Softener	Calc	09	09	40	09	09	40	09	09	0
33		В	0A	0A	OA	OA	OA	OA	VΟ	VO	none
Table 3		iv (dVg)	0.45	0.45	0.65	0.65	0.65	0.55	0.55	0.55	0.54
	Copolymer	Туре	95/5 IOA/PSMac	95/5 IOA/PSMac	90/10 IOA/PSMac	90/10 IOA/PSMac	90/10 IOA/PSMac	85/15 IOA/PSMac	85/15 IOA/PSMac	85/15 IOA/PSMac	54/36/10 IOA/HEA/PSMac
	Example Number		53	54	55	99	57	58	. 65	09	C7

		Table 3	3				
Example	Copolymer			Softener		Wet Coating Thickness (mil/µM)	J-value (X 10 ⁻⁵ cm ² /dyne)
	Type	vi (dVg)	A	Calc	Actual		
61	54/36/10 IOA/HEA/PSMac	0.54	VO	10	9.1	186/51	0.83
62	54/36/10 IOA/HEA/PSMac	0.54	0 A	20	18.3	18/381	1.18
63	54/36/10 IOA/HEA/PSMac	0.54	0 A	30	28.1	18/381	1.63
2	54/36/10 IOA/HEA/PSMac	0.54	Φ0	40	37.6	18/381	2.32
65	54/36/10 IOA/HEA/PSMac	0.54	ΨO	09	6'95	186/31	₂ 061
99	54/36/10 IOA/HEA/PSMac	0.54	0A	09	•	4/102	2302
19	95/5 IOA/PSMac	0.45	ΨO	47	45.5	18/381	40.5²
89	90/10 IOA/PSMac	0.65	0A	47	48	18/381	3.34
69	90/10 IOA/PSMac	0.65	V O	53	53.5	18/381	6.26³

		Table 3	3				
Example Number	Copolymer			Softener		Wet Coating Thickness (mil/μΜ)	J-value (X 10 ⁻⁵ cm ² /dyne)
	Туре	iv (dVg)	Ð	Calc	Actual		
70	90/10 IOA/PSMac	0.65	OA	53	,	4/102	4.432
11	85/15 IOA/PSMac	0.55	VO	47	42.2	15/381	15.13
72	85/15 IOA/PSMac	0.55	νo	53	50.7	15/381	27.03
73	57/38/5 IOA/HEA/PMIMAMac*	0.53	IPM	20	19.2	15/381	2.34³
74	57/38/5 IOA/HEA/PMMAMac*	0.53	IPM	40	39.3	15/381	34.4
75	54/36/10 IOA/HEA/PMMAMac*	0.46	IPM	20	19.6	15/381	0.79
76	54/36/10 IOA/HEA/PMMAMac*	0.46	IPM	40	38.5	15/381	93.32
80	51/34/15 IOA/HEA/PMMAMac*	0.35	None	0	0	15/381	0.42
77	51/34/15 IOA/HEA/PMMAMac*	0.35	IPM	10	9.6	15/381	0.83³

		Table 3					
Example Number	Copolymer			Softener		Wet Coating Thickness (mil/µM)	J-value (X 10 ⁻³ cm ² /dyne)
	Type	vi (dVg)	A	Calc	Actual		
78	51/34/15 10A/HEA/PMMAMac*	0.35	IPM	20	18.7	186/51	1.18³
79	51/34/15 IOA/HEA/PMMAMac*	0.35	IPM	30	27.2	186/51	1.52³
80	51/34/15 IOA/HEA/PMIMAMac*	0.35	IPM	40	36.6	18/381	334²
81	51/34/15 IOA/HEA/PMIMAMac*	0.35	пМ	90	42.1	4/102	4.46³
82	\$1/34/15 IOA/HEA/PMIMAMac*	0.35	IPM	09	45.2	4/102	4.26³
83	51/34/15 IOA/HEA/PMMAMac*	0.35	Φ0	10	6.7	18/381	0.61³
84	51/34/15 IOA/HEA/PMMAMac*	0.35	Ψ0	20	19.3	182/31	0.94³
88	\$1/34/15 IOA/HEA/PMMAMac*	0.35	٧o	30	30.5	18/381	1.22³
98	51/34/15 IOA/HEA/PMMAMac*	0.35	0 A	40	40.3	18/381	1.77³

	J-value (X 10° ³ cm²/dyne)		2.43³	3.693	4.032	9.86	36.3³	47.2	2.87²	2.99	3.62³
	Wet Coating Thickness (mil/µM)		15/381	15/381	4/102	15/381	4/102	15/381	4/102	18/381	4/102
		Actual	48.7	58.6	60.1	46.3		52.3		46	•
	Softener	Calc	20	09	09	47	47	53	53	47	47
63		Д	VO V	V	V O	OA	OA	OA	04	VO	ΨO
Table 3		i، (و/لة)	0.35	0.35	0.35	0.65	0.65	0.65	0.65	0.56	0.56
	Copolymer	Туре	51/34/15 IOA/HEA/PMMAMac*	\$1/34/15 IOA/HEA/PMMAMac*	51/34/15 IOA/HEA/PMMAMac*	57/38/5 IOA/HEA/PSMac	57/38/5 IOA/HEA/PSMac	57/38/5 IOA/HEA/PSMac	57/38/5 1OA/HEA/PSMac	54/36/10 IOA/HEA/PSMac	54/36/10 IOA/HEA/PSMac
	Example Number		87	88	89	06	91	92	93	94	95

		lable 3	~				
Example Number	Copolymer			Softener		Wet Coating Thickness (mil/µM)	J-value (X 10 ⁻⁵ cm ² /dyne)
<u></u>	Type	iv (dVg)	А	Calc	Actual		
96	54/36/10 IOA/HEA/PSMac	0.56	OA	53	51	15/381	19.1
97	54/36/10 IOA/HEA/PSMac	0.56	ΨO	53	•	4/102	125³
ప	\$1/34/15 IOA/HEA/PSMac	0.52	none	0	0	18/381	0.36
86	\$1/34/15 IOA/HEA/PSMac	0.52	OA	10	10	18/381	0.50
66	\$1/34/15 IOA/HEA/PSMac	0.52	OA	20	1.61	15/381	0.56
100	51/34/15 IOA/HEA/PSMac	0.52	0.4	30	30.4	186/51	0.77³
101	51/34/15 IOA/HEA/PSMac	0.52	0 A	40	40.5	182/381	1.16
102	51/34/15 IOA/HEA/PSMac	0.52	0 A	47	48.1	15/381	1.56
103	51/34/15 IOA/HEA/PSMac	0.52	0A	47	•	4/102	1.813

			. T	т	T -
	J-value (X 10 ⁻⁵ cm ² /dyne)		33.7	4.042	1472
	Wet Coating Thickness (mil/µM)		15/381	4/102	15/381
	L	Actual	53.9	•	61
	Softener	Calc	53	53	09
3		О	OA	VO	OA
Table 3		vi (g/lb)	0.52	0.52	0.52
	Copolymer	Type	51/34/15 IOA/HEA/PSMac	51/34/15 IOA/HEA/PSMac	51/34/15 IOA/HEA/PSMac
	Example		104	105	106

¹Average of four determinations

²Single determination

³Average of two determinations

PMMAMac* is ELVACITE 1020

Examples 107 - 129

Using the general method of Example 39, a series of coated sheet materials in which the copolymer, softener and amount of softener were varied was prepared. The copolymer, identity and amount (weight percent) of softener, wet coating thickness, and the compliance values are shown in Table 4. Unless otherwise indicated, each J-value is the average of two independent determinations. When the compliance was "not run", the formulation was too soft to be tested.

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	J-value (X 10°5	cm /dyne		0.80	1.50	2.62	4.58	64.2².	not run	not run	0.44
	Wet Coating Thickness	(Tarri amir)		18/381	15/381	15/381	18/381	4/102	4/102	4/102	18/381
	Softener			none	10% IPM*	20% IPM*	30% IPM ⁴	40% IPM ⁴	50% IPM	60% IPM	none
Table 4		.≥	(d//b)	0.54	0.54	0.54	0.54	0.54	0.54	0.54	0.50
	Copolymer	Туре		57/38/5 IOA/HEA/PMMAMac*	57/38/5 IOA/HEA/PMMAMac*	57/38/5 IOA/HEA/PMIMAMac*	57/38/5 IOA/HEA/PMMAMac*	57/38/5 IOA/HEA/PMMAMac*	57/38/5 IOA/HEA/PMIMAMac*	57/38/15 IOA/HEA/PMMAMac*	54/36/10 IOA/HEA/PMMAMac*
	Example Number			C10	107	108	109	110	111	112	C11

		Table 4			
Example	Copolymer		Softener	Wet Coating Thickness (mil/μM)	J-value (X 10 ⁻³ cm²/dyne
	Type	vi (8/lb)			
113	54/36/10 IOA/HEA/PMMAMac*	0.50	10% IPM⁴	15/381	0.69
114	54/36/10 IOA/HEA/PMMAMac*	0.50	20% IPM	18/381	0.94³
115	54/36/10 IOA/HEA/PMMAMac*	0.50	30% IPM	15/381	1.46
116	54/36/10 IOA/HEA/PMMAMac*	0.50	40% IPM	4/102	not run
117	54/36/10 IOA/HEA/PMMAMac*	0.50	80% IPM	4/102	not run
118	57/38/5 IOA/HEA/PMMAMac*	0.54	10% OA	18/381	1.63
119	57/38/5 JOA/HEA/PMMAMac*	0.54	20% OA	18/381	2.70
120	57/38/5 IOA/HEA/PMMAMac*	0.54	30% OA	15/381	4.19
121	57/38/5 IOA/HEA/PMMAMac*	0.54	40% OA	4/102	6.01

	J-value (X 10 ⁻⁵		8.27	11.8	0.60	0.89	1.19	1.56	2.65	3.99
	Wet Coating Thickness (mi/uM)		4/102	4/102	15/381	15/381	18/381	4/102	4/102	4/102
	Softener		50% OA	60% OA	10% OA	20% OA	30% OA	40% OA	50% OA	60% OA
Table 4		i (dl/g)	0.54	0.54	0.50	0.50	0.50	0.50	0.50	0.50
	Copolymer	Type	57/38/5 IOA/HEA/PMIMAMac*	57/38/5 IOA/HEA/PMIMAMac*	54/36/10 IOA/HEA/PMMAMac*					
	Example Number		122	123	124	125	126	127	128	129

PMMAMac* is ELVACITE 1020

Average of four determinations ²Single determination

³Average of three determinations

'IPM content confirmed using the test method described above.

Example 130

Copolymer (6.7306 g of 63/27/10 IOA/DMACM/PMMAMac, 47.8% solids in 95/5 w/w ethyl acetate/isopropanol, iv = 0.39 dl/g), levonorgestrel (0.0502 g) and methyl laurate (1.7606 g) were combined in an 11 dram (40.7 mL) glass vial. The vial was capped then shaken overnight on a platform shaker. The resulting 5 formulation was knife coated at a thickness of 16 mil (406 µm) onto a release liner (Daubert 164Z 5 mil [127 µM] PESTER). The coated release liner was oven dried for 4 minutes at 125°F (52°C), for 2 minutes at 185°F (85°C) and for 2 minutes at 225°F (107°C). The resulting adhesive coating contained 64.0 percent 63/27/10 IOA/HEA/PMMAMac copolymer, 1.0 percent levonorgestrel and 35.0 percent methyl laurate. The coated liner was then laminated onto the corona treated surface of a 3 mil (76.2 µm) polyethylene backing. The compliance was measured using the test method described above and found to be 4.4 X 10⁻⁵ cm²/dyne.

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Examples 131 - 178

Using the general method of Example 130, a number of coated sheet materials were prepared in order to assess the effect of increasing the amount of skin penetration enhancer(s) on the compliance of certain formulations containing levonorgestrel. The compliance was measured using the test method described above. The formulations and the J-values are shown in Table 5, where amounts are percent by weight. Except as noted, the polymethylmethacrylate macromonomer was ELVACITE 1010. PMMAMac* indicates that the polymethylmethacrylate was ELVACITE 1020.

Table 5	LN GM DDA Additional J-Value L O Enhancer(s) (cm²/dyne)	>i	1.0 0 0 30.3 ML 2.4 × 10 ⁻³	1.0 0 0 24.8 ML 2.1 x 10 ⁻⁵ AMAMac	// AAMac 1.0 0 0 17.1 DGME off scale 17.4 LG	MAMac 15.2 DGME 15.4 x 10 ⁻⁵	MAMac 1.0 0 0 12.6 DGME 5.2 x 10 ⁻⁵
			0	0	0	0	0
			0	0	0	0	0
\$	N C		1.0	1.0	1.0	1.0	1.0
Table		.≥					
	Adhesive	Туре	63/27/10 IOA/DMACM/PMMAMac	63/27/10 IOA/DMACM/PMMAMac	55/40/5 IOA/HEA/PMMAMac	55/40/5 IOA/HEA/PMIMAMac	55/40/5 IOA/HEA/PMMAMac
		Amount	68.7	74.2	64.5	68.7	74.0
1				132	133	134	135

			Table 5	le 5				
Ä		Adhesive		C.N	M _D	DDA	Additional	J-Value
o Z					1	0	Enhancer(s)	(cm²/dyne)
	Amount	Туре	.≥					•
136	78.9	55/40/5		1.0	0	0	10.1 DGME	5.0 x 10 ⁻⁵
		IOA/HEA/PMMAMac					10.0 LG	
137	65.7	55/40/5	0.51	1.0	5.0	3.0	12.8 DGME	2.6 x 10 ⁻⁵
		IOA/HEA/PMMAMac					12.5 LG	
138	6.09	55/40/5	0.51	1.0	5.0	3.0	15.0 DGME	2.9 x 10 ⁻⁵
		IOA/HEA/PMMAMac					15.1 LG	
139	55.8	55/40/5	0.51	1.0	5.0	3.0	17.6 DGME	3.4 x 10 ⁻⁵
		IOA/HEA/PMMAMac	·				17.6 LG	
140	51.1	55/40/5	15.0	1.0	5.0	3.0	20.0 DGME	8.1 x 10°5
		IOA/HEA/PMMAMac					19.9 LG	
141	65.4	55/35/10	0.42	1.0	4.9	3.1	12.7 DGME	2.2 × 10 ⁻⁵
		IOA/HEA/PMMAMac					12.9 LG	

			Table 5	S &				
Ex		Adhesive		LN	ВМ	PDA	Additional	J-Value
Š.					J	0	Enhancer(s)	(cm²/dyne)
	Amount	Туре	.≥					
142	60.5	55/35/10	0.42	1.0	4.9	3.0	15.4 DGME	1.9 x 10 ⁻⁵
		IOA/HEA/PMMAMac					15.2 LG	
143	55.7	01/5£/55	0.42	1.0	5.2	3.0	17.6 DGME	2.2 × 10 ⁻⁵
		IOA/HEA/PMMAMac					17.5 LG	
144	50.7	55/35/10	0.42	1.1	5.0	2.9	20.0 DGME	2.8 × 10 ⁻⁵
		IOA/HEA/PMMAMac					20.3 LG	
145	65.4	55/35/10	0.46	1.0	4.9	3.0	13.1 DGME	1.5 × 10 ⁻⁵
		IOA/HEA/PMMAMac*					12.6 LG	
146	60.7	95/35/10	0.46	1.1	5.4	3.0	15.0 DGME	1.8 x 10 ⁻⁵
		IOA/HEA/PMMAMac*				;	14.8 LG	
147	56.0	01/56/55	0.46	1.0	5.0	3.0	17.5 DGME	2.2 × 10 ⁻⁵
		IOA/HEA/PMMAMac•					17.5 LG	

			Tab	Table 5				
No.		Adhesive		3	GM	DDA 0	Additional Enhancer(s)	J-Value
	Amount	Type	.≥				(2)	(cm /dyne)
148	50.7	55/35/10 IOA/HEA/PMMAMac*	0.46	1.1	5.0	3.0	20.0 DGME 20.2 LG	2.4 x 10 ⁻⁵
149	52.9	63/27/10 IOA/DMACM/PMMAMac	0.48	1.0	5.1	1.0	40.0 ML	17.4 x 10 ⁻⁵
150	58.0	63/27/10 IOA/DMACM/PMMAMac	0.48	1.0	5.1	0.1	34.9 ML	9.5 x 10 ⁻⁵
151	63.1	63/27/10 IOA/DMACM/PMMAMac	0.48	0.1	5.0	1.0	29.9 ML	4.0 × 10 ⁻⁵
152	67.8	63/27/10 IOA/DMACM/PMMAMac	0.48	1.0	5.1	11	25.0 ML	3.7 x 10 ⁻⁵
153	72.9	63/27/10 IOA/DMACM/PMMAMac	0.48	1.0	5.0	1.0	20.1 ML	2.2 x 10 ⁻³
			1			1		

			Table 5	Š				
No.		Adhesive		Z,	CM	DDA 0	Additional Enhancer(s)	J-Value (cm²/dyne)
	Amount	Type	.≥1					
154	70.6	55/40/5 IOA/HEA/PMMAMac	0.51	1.0	5.0	3.0	10.3 PG 10.1 ML	3.3 × 10 ⁻⁵
155	65.0	55/40/5 IOA/HEA/PMIMAMac	0.51	1.0	5.1	3.0	12.3 PG 13.6 ML	3.1 × 10 ⁻⁵
156	60.5	55/40/5 IOA/HEA/PMMAMac	0.51	1.0	5.0	3.1	15.3 PG 15.1 ML	4.9 × 10 ⁻³
157	55.7	55/40/5 IOA/HEA/PMMAMac	0.51	1.0	5.1	3.0	17.7 PG 17.5 ML	5.3 × 10 ⁻⁵
158	51.0	55/40/5 IOA/HEA/PMMAMac	0.51	1.0	5.0	3.0	20.2 PG 19.8 ML	3.4 × 10 ⁻⁵
159	69.8	55/35/10 IOA/HEA/PMMAMac	0.42	1.0	5.2	3.0	10.0 PG 11.0 ML	1.4 x 10 ⁻⁵

			Table 5	\$ 5				
Ä		Adhesive		LN	ВМ	DDA	Additional	J-Value
No.					ר	0	Enhancer(s)	(cm²/dyne)
	Amount	Туре	'n				•	
160	66.1	55/35/10 IOA/HEA/PMMAMac	0.42	1.0	4.9	3.0	12.3 PG 12.7 ML	1.4 x 10 ⁻⁵
161	60.7	55/35/10 IOA/HEA/PMMAMac	0.42	1.0	5.0	3.0	15.3 PG 15.0 ML	2.0 × 10 ⁻⁵
162	55.8	55/35/10 IOA/HEA/PMMAMac	0.42	1.0	5.0	3.0	17.7 PG 17.5 ML	2.3 x 10 ⁻⁵
163	50.7	55/35/10 IOA/HEA/PMIMAMac	0.42	1.0	5.3	3.0	20.2 PG 19.8 ML	2.7 x 10 ⁻⁵
164	72.0	60/15/15/10 IOA/DMACM/HEA/PMMAMac	0.47	1.0	5.0	2.0	14.3 ML 5.7 DIPA	2.0 x 10 ⁻⁵
165	67.3	60/15/10 IOA/DMACM/HEA/PMMAMac	0.47	1.0	5.0	2.1	17.8 ML 6.8 DIPA	2.4 x 10 ⁻⁵

	J-Value	(cm²/dyne)		5.0 × 10 ⁻⁵		7.8 × 10 ⁻⁵		16.6 × 10 ⁻⁵		15.4 × 10 ⁻⁵		24.8 x 10 ⁻⁵		1.8 x 10 ⁻⁵	
	Additional	Enhancer(s)		21.8 ML	8.4 DIPA	25.4 ML	9.6 DIPA	28.8 ML	11.0 DIPA	20.3 ML		24.9 ML		20.9 ML	
	DDA	0	<u></u>	2.1		2.0		2.0		1.0		1.1		1.0	
	MS	1		5.0		5.1		5.2		5.0		5.0		4.9	
e 5	LN			1.0		1.0		1.0		1.0		1.0		1.0	
Table 5			'n	0.47		0.47	:	0.47		0.47		0.47		0.53	
	Adhesive		Туре	60/15/15/10	IOA/DMACM/HEA/PMMAMac	60/15/15/10	IOA/DMACM/HEA/PMMAMac	01/51/5/109	IOA/DMACM/HEA/PMMAMac	68/27/5	IOA/DMACM/PMMAMac	68/27/5	IOA/DMACM/PMMAMac	50/40/10	IOA/DMACM/PMMAMac
		i	Amount	61.7		6.95		52.0	_	72.7		0.89		72.2	
	Ex	o Z		166		167		168		169		170		171	

			Tab	Table 5				
Ex No.		Adhesive		LN	GM L	DDA 0	Additional Enhancer(s)	J-Value
	Amount	Type	.≥					(cur /dyne)
172	67.7	50/40/10 IOA/DMACM/PMMAMac	0.53	1.0	5.0	1.0	25.3 ML	2.7 x 10 ⁻³
173	63.5	50/40/10 IOA/DMACM/PMMAMac	0.53	1.0	4.9	1.0	29.6 ML	5.2 x 10 ⁻⁵
174	58.3	50/40/10 IOA/DMACM/PMMAMac	0.53	1.1	5.0	1.1	34.5 ML	10.7 x 10 ⁻⁵
175	53.0	50/40/10 IOA/DMACM/PMMAMac	0.53	1.0	5.1	1:1	39.8 ML	21.5 x 10°5
176	71.0	65/15/15/5 IOA/DMACM/HEA/PMMAMac	0.47	1.0	5.0	2.0	13.7 ML 7.3 DIPA	8.8 × 10 ⁻⁵
177	66.7	65/15/15/5 IOA/DMACM/HEA/PMIMAMac	0.47	1.0	5.1	2.0	17.5 ML 7.7 DIPA	13.2 x 10 ⁻⁵

			Table 5	5 5				
益		Adhesive		Ľ	Βğ	DDA	LN GM DDA Additional	J-Value
Š.						0	Enhancer(s)	(cm²/dyne)
	Amount	Type	.≥					
178	62.6	65/15/15/5	0.47	1.0 5.1 2.0	5.1	2.0	20.3 ML	22.9 × 10 ⁻⁵
		IOA/DMACM/HEA/PMMAMac	-				9.0 DIPA	

In Vitro Skin Penetration Test Method

The skin penetration data given in the examples below was obtained using the following test method. A Diffusion cell 20 of the type shown in Figure 1 is used. Human cadaver skin (Dermatomed skin about 500µM thick obtained from a skin bank) is used. A shown in Figure 2, the skin 22 is mounted epidermal side up between upper portion 24 and lower portion 26 of the cell, which are held together by means of ball joint clamp 28.

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The portion of the cell below the mounted skin is completely filled with receptor fluid (30% N-methyl-2-pyrrolidone in water) such that the receptor fluid is in contact with the skin. The receptor fluid is stirred using a magnetic stirrer (not illustrated). The sampling port 30 is covered except when in use.

When a transdermal delivery device is evaluated, the skin is placed across the orifice of the lower portion of the diffusion cell, the release liner is removed from a 2.0 cm² patch and the patch is applied to the skin and pressed to cause uniform contact with the skin. The diffusion cell is assembled and the lower portion is filled with 10 mL of warm (32°C) receptor fluid.

The cell is the placed in a constant temperature (32 ± 2°C) and humidity (50 ± 10% relative humidity) chamber. The receptor fluid is stirred by means of a magnetic stirrer throughout the experiment to assure a uniform sample and a reduced diffusion barrier on the dermal side of the skin. The entire volume of receptor fluid is withdrawn at specified time intervals (6, 12, 24, 48 and 72 hours) and immediately replaced with fresh fluid. The withdrawn fluid is filtered through a 0.45 μM filter. A 1 mL portion of filtrate is then analyzed for levonorgestrel using high performance liquid chromatography (Column: 15 cm X 4.6 mm I.D. ZORBAXTM RX-C18 from DuPont, 5 μM particle size; Mobile Phase: 60/40 v/v water/acetonitrile; Flow Rate: 1.5 mL/min; Run Time: 11.0 min; Detection: uv at 230 nm). The cumulative amount of levonorgestrel penetrating the skin is calculated. The greatest slope of a plot of the cumulative penetration versus time is reported as steady state levonorgestrel flux measured in μg/cm²/hour.

Example 179

Levonorgestrel (19.85 g), methyl laurate (330.8 g), propylene glycol (198.5 g), glyceryl monolaurate (33.08 g), N,N-dimethyldodecylamine-N-oxide (19.85 g) and copolymer (1803 g of 55/40/5 IOA/HEA/PMMAMac copolymer, 40% solids in 95/5 w/w ethyl acetate/isopropanol, which had been dried then resolvated, iv = 0.59 dl/g after drying) were placed in a 1 gallon (3.8 L) high density polyethylene carboy. The carboy was tightly capped then placed on a roller/shaker for 19 hours. The carboy was allowed to stand until all entrapped air bubbles had dissipated. The resulting formulation was knife coated at a wet thickness of 16 mil (406 µM) onto a silicone coated polyester (5 mil, 127 µM) film. The coated release liner was oven dried at 127°F (53°C) for 30 minutes. The resulting adhesive coating contained 1.5 percent levonorgestrel, 15.0 percent propylene glycol, 25.0 percent methyl laurate, 2.5 percent glyceryl monolaurate, 1.5 percent N,N-dimethyldodecylamine-N-oxide, and 54.5 percent 55/40/5 IOA/HEA/PMMAMac copolymer. The coated liner was allowed to cool for 10 minutes then it was laminated to the corona treated side of a 2 mil (51 µM) polypropylene film. The compliance was measured using the test method described above and found to be 6.57 X 10⁻⁵ cm²/dynes. Skin penetration through human cadaver skin was measured using the test method described above; the steady state flux was found to be 0.166 µg/cm²/hr.

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Example 180

Levonorgestrel (18.29 g), methyl laurate (457.2 g), glyceryl monolaurate (65.31 g), N,N- dimethyldodecylamine-N-oxide (13.06 g) and copolymer (1401 g of 50/40/10 IOA/DMACM/PMMAMac copolymer, 53.7% solids in 95/5 w/w ethyl acetate/isopropanol, which had been dried then resolvated, iv = 0.55 dl/g before drying; iv = 0.52 dl/g after drying) were placed in a 1 gallon (3.8 L) high density polyethylene carboy. The carboy was tightly capped then placed on a roller/shaker for 19 hours. The carboy was allowed to stand until all entrapped air bubbles had dissipated. The resulting formulation was knife coated at a wet thickness of 12 mil (305 μ M) onto a silicone coated polyester (5 mil, 127 μ M) film. The coated release liner was oven dried at 127°F (53°C) for 80 minutes. The resulting adhesive

coating contained 1.4 percent levonorgestrel, 35.0 percent methyl laurate, 5.0 percent glyceryl monolaurate, 1.0 percent N,N-dimethyldodecylamine-N-oxide, and 57.6 percent 50/40/10 IOA/DMACM/PMMAMac copolymer. The coated liner was allowed to cool for 10 minutes then it was laminated to the corona treated side of a 2 mil (51 µM) polypropylene film. The compliance was measured using the test method described above and found to be 5.74 X 10⁻⁵ cm²/dynes. Skin penetration through human cadaver skin was measured using the test method described above; the steady state flux was found to be 0.148 µg/cm²/hr.

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Example 181

Levonorgestrel (18.04 g), methyl laurate (264.6 g), tetraglycol (96.23 g), glyceryl monolaurate (60.14 g), N,N-dimethyldodecylamine-N-oxide (12.03 g) and copolymer (1400 g of 50/40/10 IOA/DMACM/PMMAMac copolymer, 53.7% solids in 95/5 w/w ethyl acetate/isopropanol, which had been dried then resolvated, iv = 0.55 dl/g before drying; iv = 0.52 dl/g after drying) were placed in a 1 gallon (3.8 L) high density polyethylene carboy. The carboy was tightly capped then placed on a roller/shaker for 19 hours. The carboy was allowed to stand until all entrapped air bubbles had dissipated. The resulting formulation was knife coated at a wet thickness of 13 mil (330 µM) onto a silicone coated polyester (5 mil, 127 μM) film. The coated release liner was oven dried at 127°F (53°C) for 75 minutes. The resulting adhesive coating contained 1.5 percent levonorgestrel, 22.0 percent methyl laurate, 8.0 percent tetraglycol, 5.0 percent glyceryl monolaurate, 1.0 percent N,N-dimethyldodecylamine-N-oxide, and 62.5 percent 50/40/10 IOA/DMACM/PMMAMac copolymer. The coated liner was allowed to cool for 10 minutes then it was laminated to the corona treated side of a 2 mil (51 μ M) polypropylene film. The compliance was measured using the test method described above and found to be 8.72 X 10⁻⁵ cm²/dynes. Skin penetration through human cadaver skin was measured using the test method described above; the steady state flux was found to be 0.131 µg/cm²/hr.

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Example 182

Copolymer (50.13 g of 57/38/5 IOA/HEA/PMMAMac, 39.5% solids in 97/3 ethyl acetate/isopropanol, iv = 0.69 dl/g) and nicotine (5.04 g) were combined in a glass jar. The jar was capped and shaken for 15 minutes. The resulting formulation was knife coated at a wet thickness of 8 mil (203 μM) onto a silicone coated polyester release liner (5 mil (127 μM) Daubert). The coated release liner was oven dried at 110°F (43°C) for 30 minutes. The resulting coating theoretically contained 79.71 percent 57/38/5 IOA/HEA/PMMAMac copolymer and 20.29 percent nicotine. The coated liner was laminated to a backing (1109 SCOTCHPAKTM tan, polyester film laminate, available from the 3M Company). The compliance was measured 4 hours after the laminate was prepared using the test method described above and found to be 1.79 X 10⁻⁵ cm²/dyne. The compliance was measured again after the laminate had sat overnight and was found to be 1.5 X 10⁻⁵ cm²/dyne (average of two independent determinations).

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Example 183

The formulation prepared in Example 182 was knife coated at a wet thickness of 6 mil (152 μM) onto a silicone coated polyester release liner (5 mil (127 μM) Daubert). The coated release liner was allowed to dry at ambient temperature (22°C) for 100 minutes. The resulting coating theoretically contained 79.71 percent 57/38/5 IOA/HEA/PMMAMac copolymer and 20.29 percent nicotine. The coated liner was laminated to a backing (1109 SCOTCHPAKTM tan, polyester film laminate, available from the 3M Company). The compliance was measured after the laminate had sat over the weekend and was found to be 2.4 X 10⁻⁵ cm²/dyne (average of two determinations).

Example 184

Copolymer (10.0 g of 55/9/28/8 2-ethylhexylacrylate/vinyl acetate/tetrahydrofurfuryl acrylate/ELVACITE 1020 PMMAMac 37.28 % solids in 90/10 w/w ethyl acetate/isopropanol, iv = 0.706 dl/g) and isopropyl myristate (0.93 g) were combined then mixed to provide a homogeneous formulation. The

formulation was coated at a wet thickness of 15 mil (381 μ M) onto a polyethylene terephthalate film then air dried to provide a pressure sensitive adhesive with clean release from skin.

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Example 185

Copolymer (10.0 g of 55/9/28/8 2-ethylhexylacrylate/vinyl acetate/tetrahydrofurfuryl acrylate/ELVACITE 1020 PMMAMac 37.28 % solids in 90/10 w/w ethyl acetate/isopropanol, 0.706 dl/g) and isopropyl myristate (1.60 g) were combined then mixed to provide a homogeneous formulation. The formulation was coated at a wet thickness of 15 mil (381 μ M) onto a polyethylene terephthalate film then air dried to provide a pressure sensitive adhesive with clean release from skin.

Example 186

15 Copolymer (10.0 g of 82/10/8 IOA/2-hydroxyethyl methacrylate/ELVACITE 1020 PMMAMac 38.7% solids in 95/5 w/w ethyl acetate/isopropanol, iv = 0.378 dl/g) and oleyl alcohol (0.97 g) were combined then mixed to provide a homogeneous formulation. The formulation was coated at a wet thickness of 15 mil (381 μM) onto a polyethylene terephthalate film then air dried to provide a pressure sensitive adhesive with clean release from skin.

Example 187

Copolymer (10.0 g of 77/4/15/4 IOA/acrylamide/DMACM/ELVACITE 1020 PMMAMac 39.5% solids in 95/5 w/w ethyl acetate/isopropanol, iv = 0.443 dl/g) and isopropyl myristate (0.99 g) were combined then mixed to provide a homogeneous formulation. The formulation was coated at a wet thickness of 15 mil (381 μ M) onto a polyethylene terephthalate film then air dried to provide an aggressive pressure sensitive adhesive with clean release from skin.

Example 188

Copolymer (10.0 g of 74/9/9/8 2-ethylhexyl acrylate/N-vinyl pyrrolidone/2-hydroxyethyl acrylate/ELVACITE 1020 PMMAMac 39.4% solids in 95/5 w/w ethyl acetate/isopropanol, iv = 0.365 dl/g) and isopropyl myristate (0.99 g) were combined then mixed to provide a homogeneous formulation. The formulation was coated at a wet thickness of 15 mil (381 μ M) onto a polyethylene terephthalate film then air dried to provide an aggressive pressure sensitive adhesive with clean release from skin.

10 Example 189

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Copolymer (10.0 g of 55/9/28/8 IOA/butyl methacrylate/ethoxy ethoxy ethyl acrylate/ELVACITE 1020 PMMAMac 38.3% solids in 95/5 w/w ethyl acetate/isopropanol, iv = 0.78 dl/g) and oleyl alcohol (0.96 g) were combined then mixed to provide a homogeneous formulation. The formulation was coated at a wet thickness of 15 mil (381 μ M) onto a polyethylene terephthalate film then air dried to provide a pressure sensitive adhesive with clean release from skin.

Example 190

Copolymer (10.0 g of 55/9/28/8 IOA/butyl methacrylate/ethoxy ethoxy ethyl acrylate/ELVACITE 1020 PMMAMac 38.3% solids in 95/5 w/w ethyl acetate/isopropanol, iv = 0.78 dl/g) and oleyl alcohol (1.64 g) were combined then mixed to provide a homogeneous formulation. The formulation was coated at a wet thickness of 15 mil (381 μM) onto a polyethylene terephthalate film then air dried to provide a pressure sensitive adhesive with limited tack and with clean release from skin.

Example 191

Copolymer (10.0 g of 55/9/28/8 IOA/butyl acrylate/ethoxy ethoxy ethyl acrylate/ELVACITE 1020 PMMAMac 38.5% solids in 95/5 w/w ethyl acetate/isopropanol, iv = 0.78 dl/g) and oleyl alcohol (0.96 g) were combined then mixed to provide a homogeneous formulation. The formulation was coated at a wet thickness of 15 mil (381 μ M) onto a polyethylene terephthalate film then air dried to provide a pressure sensitive adhesive with clean release from skin.

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Example 192

10 Copolymer (10.0 g of 55/9/28/8 IOA/butyl acrylate/ethoxy ethoxy ethyl acrylate/ELVACITE 1020 PMMAMac 38.5% solids in 95/5 w/w ethyl acetate/isopropanol, iv = 0.78 dl/g) and oleyl alcohol (1.65 g) were combined then mixed to provide a homogeneous formulation. The formulation was coated at a wet thickness of 15 mil (381 μ M) onto a polyethylene terephthalate film then air dried to provide a pressure sensitive adhesive with limited tack and with clean 15 release from skin.

Example 193

Copolymer (100 g of 61/37/2 IOA/VoAc/PSMac, 34 percent solids in 84/16 ethyl acetate/toluene, iv = 0.87 dl/g) and oleyl alcohol (14.57 g) were combined in a 20 glass jar. The jar was placed on a roller mixer overnight. The resulting formulation was knife coated at a wet thickness of about 7 mil (178 μ M) onto a 2 mil (51 µM) polyethylene terephthalate film. The coated film was oven dried at 110°F (43°C) for 20 minutes. The resulting coating theoretically contained 70 percent 61/37/2 IOA/VoAc/PSMac copolymer and 30 percent oleyl alcohol. The coated film was folded back onto itself to form a "sandwich" and the compliance was measured using the test method described above. The compliance was found to be 6.8 X 10⁻⁵ cm²/dyne (average of three independent determinations).

Examples 194 - 218

Using the general method of Example 193, a series of coated sheet materials in which the copolymer, softener and amount of softener were varied was prepared. The copolymer, identity and amount (weight percent) of softener, and the compliance values are shown in Table 6 where each J-value is the average of three independent determinations. The polymethylmethacrylate macromonomer used was ELVACITE 1020.

Table 6		J- value	(X 10 ⁻⁵ cm ² /dyne)	Type iv (dl/g)	61/37/2 IOA/VoAc/PSMac 0.87	200% IDM	30% TBM	100 TOUR	0.87		1.02 none 0.65	61/37/2 IOA/VoAc/PSMac 1.02 20% IPM 8.3	-	1 02 400, 1104	1 02 2000 O	102 A200 A200 A200 A200 A200 A200 A200 A			20,5 15 TO VOROT SIMBC 0.89 20% IPM 2.3	58/37/5 IOA/VoAc/PSMac 0.89 2002, my
	Example	Number			C12	194	561	196	197	C13	100	130	661	200	201	202	C14	203	700	704

	ener J- value	(X 10 ⁻⁵ cm ² /dyne)		IPM >20	0A 1.1	OA >20	none 0.44	IPM 3.9	IPM 11.2	40% IPM >20	30% OA 1.6	40% OA >20	none 0.15	30% OA 0.32	none 0.16	30% OA 0.36	0.4
	Softener		iv (dl/g)	0.89 40% IPM	0.89 30% OA	0.89 40% OA	1.02 no	1.02 20% IPM	1.02 30% IPM	1.02 40%	1.02	1.02	0.815 no	0.815 30%	0.92 no	0.92 30%	1.05
Table 6	Copolymer		Туре	58/37/5 IOA/VoAc/PSMac	53/37/10 IOA/VoAc/PMMAMac	53/37/10 IOA/VoAc/PMIMAMac	53/37/10 IOA/VoAc/PMIMAMac	53/37/10 IOA/VoAc/PMIMAMac	58/37/5 IOA/VoAc/PMIMAMac								
	Example	Number		205	206	207	C15	208	500	210	211	212	C16	213	C17	214	C18

	J- value	(A 10 cm /dyne)		290	0.31	0.71	0.37	0.7	6	ø. Э
	Softener			30% OA	30% IPM		none	30% OA	30% IPM	W 17 2/25
Table 6		iv (41/n)	(8,18)	1.05	1.05		CII	1.15	1.15	
I.	Copolymer	Type	2012/EU83	30/3 //3 IOA/ VOAC/PMIMAMac	58/37/5 IOA/VoAc/PMMAMac	58/37/5 IOA/VoAc/PMMAMac		38/3//3 IOA/VOAC/PMMAMac	58/37/5 IOA/VoAc/PMMAMac	
	Example Number		215		216	C19	217		218	

Example 219

Copolymer (58/37/5 IOA/VoAc/PSMac, 34 percent solids in 84/16 ethyl acetate/toluene, iv = 0.89 dl/g) was knife coated at a wet thickness of about 7 mil (178 µM) onto a 2 mil (51 µM) polyethylene terephthalate film. The coated film was oven dried at 160°F (71°C) for 20 minutes and then at 210°F (99°C) for 10 minutes. Patches (5 cm² circles) each containing 0.044 g of dry adhesive were cut from the adhesive coated film. Nicotine (0.011 g) was placed on top of the adhesive in each patch using a micropipette to provide a patch with an adhesive layer containing 20 percent by weight of nicotine. The adhesive layer was covered with a release liner (SCOTCHPAKTM 1022) and allowed to equilibrate overnight. The rate of release of nicotine from the patch was determined using the test method described below. The results are shown in Table 7 below where each entry is the average of three independent determinations.

Example 220

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The method of Example 219 was repeated using a 58/37/5 IOA/VoAc/PSMac having an iv = 1.02 dl/g. The rate of release of nicotine from the patch was determined using the test method described below. The results are shown in Table 7 below where each entry is the average of three independent determinations.

In-vitro Release of Nicotine

This method describes the dissolution test procedure used to evaluate invitro release characteristics of nicotine transdermal delivery patches.

The method uses a Hanson Dissolution Apparatus with the dissolution media temperature set at 32°C; the paddle speed set at 50 rpm; and the paddle height above the sample set at 25 mm.

Each patch (5 cm²) is affixed with double sided adhesive tape to a separate stainless steel plate so that the release liner is facing upward (backing is in direct contact with the double sided tape). Each dissolution flask is charged with 500 mL

0.1 M phosphate buffer (pH 6.0) and the temperature of the buffer is allowed to equilibrate at 32 ± 0.5 °C.

The release liner is removed from the patch and the mounted patch is placed in the dissolution flask. At 5, 10, 20, 30, 60, 90, 120, 240, 480 and 720 minutes, 4 mL samples are withdrawn and analyzed for nicotine content using uv sprectrophotometry with the wavelength set at 262 nm using a 1 cm flow through the spectrophotometer cell. The results are reported as the cumulative percent nicotine released.

	Table 7	
	In-vitro Nicotine Release	
Time (minutes)	Cumulative Percen	t Nicotine Released
	Example 219	Example 220
0	0	0
5	36.7	38.4
10	44.2	46.6
20	55.8	60.3
30	65.9	68.7
60	77.5	80.0
90	80.5	84.6
120	84.9	87.2
240	87.6	89.3
480	88.5	90.4
720	89.8	90.9

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Example 221

Using the method of Example 219, patches having an adhesive layer containing 25 percent by weight of nicotine were prepared using a 53/37/10 IOA/VoAc/ELVACITE 1020 copolymer having an iv = 0.92 dl/g. The adhesive

layer of the patch had many air bubbles. The compliance was found to be 1.5 X 10⁻⁵ cm²/dyne (average of three independent determinations).

Example 222

Using the method of Example 219, patches having an adhesive layer containing 25 percent by weight of nicotine were prepared using a 58/37/5 IOA/VoAc/ELVACITE 1020 copolymer having an iv = 1.15 dl/g. The compliance was found to be 0.9 X 10⁻⁵ cm²/dyne (average of three independent determinations).

10 Example 223

Propylene glycol (1.52 g), methyl laurate (2.54 g), glyceryl monolaurate (0.25 g), N,N-dimethyldodecylamine-N-oxide (0.15 g), dried copolymer (5.53 g of 55/40/5 IOA/HEA/PMMAMac, iv = 0.45 dl/g prior to drying) and solvent (15 g of 95/5 w/w ethyl acetate/isopropanol) were combined and mixed to provide a homogeneous coating formulation. The formulation was coated at a wet thickness of 20 mil (508 μM) onto a silicone coated polyester release liner (Daubert PESTER). The coated release liner was oven dried for 4 minutes at 43°C, for 3 minutes at 85°C, and for 2 minutes at 107°C. The coated release liner was then laminated to the corona treated side of a clear 2 mil (51 μM) polypropylene film. Patches (circular, 5 cm²) were die cut from the resulting laminate. One patch was applied to the left forearm of a human subject. A second patch was applied to the right forearm of the same subject. The percent of patch surface adhering to skin was approximated by visual assessment through the clear backing. The results are shown in Table 8 below.

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Examples 224 - 261

Using the general method of Example 223, a number of patches were prepared and the adhesion to skin evaluated in order to assess the effect of copolymer composition, copolymer inherent viscosity, wet coating thickness, softener composition and the amount of softener on adhesion to skin. The formulations (amounts are percent by weight) and adhesion evaluations are shown

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in Table 8 below wherein the absence of an entry indicates that the adhesion was not assessed at that time point, "OFF" means that the patch fell off by itself, and "R" means that the patch was removed by the subject. All adhesion testing was conducted on the same subject and unless otherwise indicated the patch was adhered to the left forearm.

														,	
				Day 4		20		20	. —	09		50		45	
	(%			Day 3		99		85		75	_	88		85	
	Adhesion (%)			Day 2		85	-	95		06		95		8	
	ΨC			Day 0 Day 1											
:				Day 0		<u>8</u>		200		<u>8</u>		901		100	
	Wet Coating	Thickness	(miVµM)			20/208		20/208		20/208		20/208		20/208	
Table 8	Softener					15.2 PG; 25.4 ML;	2.5 GML	15.2 PG; 25.4 ML;	2.5 GML	10.1 PG; 30.5 ML;	2.5 GML	10.1 PG; 30.5 ML;	2.5 GML	5.1 PG; 35.5 ML;	2.5 GML
				Ņ	(dl/g)	0.45		0.45		0.45		0.45		0.45	
	Copolymer			Туре		55/40/5 IOA/HEA/PMIMAMac		55/40/5 IOA/HEA/PMMAMac		55/40/5 IOA/HEA/PMIMAMac		55/40/5 IOA/HEA/PMMAMac		55/40/5 IOA/HEA/PMIMAMac	
	Example	Number				2231		22412	!	2251		226 ^{1,2}		227	

_	1			1		_									
				Day 4	_	25				~		2		æ	:
	 @			Day 3		75		OFF		09		01		86~	
	Adhesion (%)	•		Day 2		8		65		86		85		8	
				Day 1	•			8		100		95	-	100	
				Day 0		100		100		100	-	100		100	
	Wet Coating	Thickness	(mil/µM)			20/208		20/208		20/508		20/508		20/508	
Table 8	Softener					5.1 PG; 35.5 ML;	2.5 GML	15.2 PG; 25.4 ML;	2.5 GML	15.2 PG; 25.4 ML;	2.5 GML	10.1 PG; 30.5 ML;	2.5 GML	10.1 PG; 30.5 ML;	2.5 GML
				.≥	(g/lb)	0.45		0.75		0.75		0.75		0.75	
	Copolymer			Type		55/40/5 IOA/HEA/PMIMAMac		60/35/5 IOA/HEA/PMMAMac		60/35/5 IOA/HEA/PMIMAMac		60/35/5 IOA/HEA/PMIMAMac		60/35/5 IOA/HEA/PMMAMac	
	Example	Number				22812		2291		230''.2		231		2321.2	

								_		_			-		
				Day 4	_			R		90	OFF			75	
-	ૼ			Day 3		R		56~		09	99		OFF	78	
	Adhesion (%)			Day 2		10		100		08	02 ·	OFF	9	08	
	Ad			Day 1		95		100		56	82	20	8	08	
				Day 0		100		100		00 1	100	<u>8</u>	8	<u>8</u>	
	Wet Coating	Thickness	(mil/µM)			20/508		20/208		15/381	18/381	15/381	15/381	18/381	
Table 8	Softener					5.1 PG; 35.5 ML;	2.5 GML	5.1 PG; 35.5 ML;	2.5 GML	30 OA	44 OA	30 ML	44 ML	10.2 PG; 30.5 ML;	2.5 GML
				.≥	(g/lb)	0.75		0.75		0.45	0.45	0.45	0.45	89.0	
	Copolymer			Type		60/35/5 IOA/HEA/PMIMAMac		60/35/5 IOA/HEA/PMMAMac		55/40/5 IOA/HEA/PMIMAMac	55/40/5 IOA/HEA/PMIMAMac	55/40/5 IOA/HEA/PMIMAMac	55/40/5 IOA/HEA/PMIMAMac	59/40/1 IOA/HEA/PMIMAMac*	
	Example	Number	-			2331		2341,2		235	236	237	238	239	_

ML 0.5 ML; 15/381 ML 0.5 ML; 25/635 ML	2.5 GML 10.2 PG; 30.5 ML; 2.5 GML 10.2 PG; 30.5 ML; 2.5 GML	2.5 GML 0.69 10.2 PG; 30.5 ML; 2.5 GML 0.68 10.2 PG; 30.5 ML; 2.5 GML
ML 0.5 ML; 25/635 ML	10.2 PG; 30.5 ML; 2.5 GML	0.63 10.2 PG; 30.5 ML; 2.5 GML
	· .	0.63
58/38/4 IOA/HEA/PMMAMac* 0.6 59/40/1 IOA/HEA/PMMAMac* 0.6 59/39/2 IOA/HEA/PMMAMac* 0.6	58/38/4 IOA/HEA/PMMAN 59/40/1 IOA/HEA/PMMAN 59/39/2 IOA/HEA/PMMAN	

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														,		
	!			Day 4		80		8				2		R		2
	(%			Day 3		~88		95		OFF		75		99	OFF	20
	Adhesion (%)			Day 2		96		96~		65		08		20	20	55
	Ad			Day 1		100		86~	_	99		85		85	70	75
				Day 0		100		100		80		95		100	95	95
	Wet Coating	Thickness	(mil/µM)		-	25/635		25/635		15/381		186/51		15/381	15/381	15/381
Table 8	Softener					10.2 PG; 30.5 ML;	2.5 GML	10.2 PG; 30.5 ML;	2.5 GML	10.2 PG; 30.5 ML;	2.5 GML	10.2 PG; 30.5 ML;	2.5 GML	44 EO	44 OA	44 ML
				.≥	(dl/g)	0.62		69.0		0.55		0.32		0.55	0.55	0.55
	Copolymer			Type		58/39/3 IOA/HEA/PMIMAMac*		58/38/4 IOA/HEA/PMMAMac*		57/38/5 IOA/HEA/PSMac		57/38/5 IOA/HEA/PSMac		57/38/5 IOA/HEA/PSMac	57/38/5 IOA/HEA/PSMac	57/38/5 IOA/HEA/PSMac
	Example	Number				2451		246		247		248		249	250	251

	,										_				
				Day 4		2			0FF			OFF	OFF	OFF	
	(%)			Day 3		75			~63			35	20	45	
	Adhesion (%)			Day 2	_	80		24	56~			35	20	45	
	¥			Day 1		95	OFF	30	86~	OFF	OFF	.20	80	20	
				Day 0		100	100	100	100	001	100	100	100	81	
	Wet Coating	Thickness	(mil/µM)			20/508	20/208	20/208	20/208	20/208	20/508	20/508	20/208	20/208	
Table 8	Softener					30 EO	30 OA	30 ML	30 IPM	44 EO	44 OA	44 ML	44 IPM	10.2 PG; 30.5 ML;	2.5 GML
				Ņ	(g/lb)	0.55	0.55	0.55	0.55	0.55	0.55	0.55	0.55	0.32	
	Copolymer		-	Туре		57/38/5 IOA/HEA/PSMac									
	Example	Number				252	253	254	255	256	257	258	259	260	

				Day 4		OFF	
	9			Day 3		OFF	
	Adhesion (%)			Day 0 Day 1 Day 2 Day 3 Day 4		80	··· -
	Adl			Day 1		80	
				Day 0		100	
	Wet Coating	Thickness	(mil/µM)			20/208	
Table 8	Softener					10.2 PG; 30.5 ML;	2.5 GML
				^!	(dl/g)	0.55	
	Copolymer			Type		57/38/5 IOA/HEA/PSMac	
	Example	Number				261	

*PMMAMac is ELVACITE 1020

¹Formulation also contained 1.5% DDAO

²Adhesion test conducted on subject's right arm

WHAT IS CLAIMED IS:

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1. A transdermal drug delivery device, comprising:

- (1) a backing;
- (2) a matrix adhered to one side of the backing and comprising
 - (a) a copolymer comprising

(i) one or more A monomers selected from the group consisting of alkyl acrylates containing 4 to 10 carbon atoms in the alkyl group and alkyl methacrylates containing 4 to 10 carbon atoms in the alkyl group; and

(ii) optionally one or more ethylenically unsaturated B monomers copolymerizable with the A monomer; and

(iii) a macromonomer copolymerizable with the A and B monomers defined above and having a molecular weight in the range 500-500,000;

(b) a softener dissolved in the copolymer; and,

(c) if the softener is not therapeutically effective, a therapeutically effective amount of a drug,

wherein the structure and amount of the comonomers in the copolymer, the inherent viscosity of the copolymer, and the amount and structure of the drug and the softener are such as to provide the matrix with a compliance value in the range 2×10^{-6} cm²/dyne to about 4×10^{-3} cm²/dyne.

- 2. A transdermal drug delivery device according to Claim 1, wherein the B monomer or monomers comprises a functional group selected from the group consisting of carboxylic acid, carboxylic acid ester, hydroxy, sulfonamide, urea, carbamate, carboxamide, amine, oxy, oxo, and cyano.
- A transdermal drug delivery device according to Claim 1, wherein the B monomer or monomers are selected from the group consisting of acrylic acid, methacrylic acid, maleic acid, a hydroxyalkyl acrylate containing 2 to 4 carbon atoms in the hydroxyalkyl group, a hydroxyalkyl methacrylate containing 2 to 4

carbon atoms in the hydroxyalkyl group, acrylamide, methacrylamide, an alkyl substituted acrylamide containing 1 to 8 carbon atoms in the alkyl group, diacetone acrylamide, a dialkyl acrylamide having 1 or 2 carbon atoms in the alkyl group, N-vinyl-N-methyl acetamide, N-vinyl valerolactam, N-vinyl caprolactam, N-vinyl-2-pyrrolidone, glycidyl methacrylate, alkoxyethyl acrylate containing 1 to 4 carbon atoms in the alkoxy group, alkoxyethyl methacrylate containing 1 to 4 carbon atoms in the alkoxy group, 2-ethoxyethoxyethyl acrylate, furfuryl methacrylate, furfuryl acrylate, tetrahydrofurfuryl acrylate, tetrahydrofurfuryl methacrylate, propylene glycol monomethacrylate, propylene glycol monoacrylate, polyethylene glycol acrylate, polyethylene glycol methacrylate, polyethylene glycol methyl ether acrylate, polyethylene oxide methyl ether acrylate, di(lower)alkylamino ethyl acrylate, di(lower)alkylamino ethyl methacrylate, di(lower)alkylaminopropyl methacrylamide, acrylonitrile, methacrylonitrile, and vinyl acetate.

- 4. A transdermal drug delivery device according to Claim 1, wherein the A monomer is present in an amount of about 40 to about 95 percent by weight, based on the total weight of all monomers in the copolymer.
- 5. A transdermal drug delivery device according to Claim 1, wherein
 20 the A monomer is present in an amount of about 50 to about 70 percent by weight,
 based on the total weight of all monomers in the copolymer.
 - 6. A transdermal drug delivery device according to Claim 1, wherein the A monomer is selected from the group consisting of isooctyl acrylate, 2-ethylhexyl acrylate, butyl acrylate, and cyclohexyl acrylate.
 - 7. A transdermal drug delivery device according to Claim I, wherein the B monomer is present in an amount from 0 to 60 percent by weight based on the total weight of the copolymer.

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8. A transdermal drug delivery device according to Claim 1, wherein the B monomer is present in an amount of greater than 25 percent by weight based on the total weight of the copolymer, to about 50 percent by weight based on the total weight of the copolymer.

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- 9. A transdermal drug delivery device according to Claim 1, wherein the B monomer is selected from the group consisting of hydroxyethyl acrylate, hydroxyethyl methacrylate, glyceryl acrylate, N,N-dimethyl acrylamide, 2-ethoxyethoxyethyl acrylate, 2-ethoxyethyl acrylate, tetrahydrofurfuryl acrylate, acrylic acid, acrylamide, and vinyl acetate.
- 10. A transdermal drug delivery device according to Claim 1, wherein the macromonomer has a molecular weight in the range 5,000-30,000.
- 15 I1. A transdermal drug delivery device according to Claim 1, wherein the macromonomer is present in an amount of not more than about 15% by weight based on the total weight of all monomers in the copolymer.
- 12. A transdermal drug delivery device according to Claim 1, wherein
 20 the macromonomer is present in an amount of not more than about 5% by weight based on the total weight of all monomers in the copolymer.
 - 13. A transdermal drug delivery device according to Claim 1, wherein the macromonomer is a compound of the formula

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wherein X is a moiety comprising an ethylenically unsaturated group copolymerizable with the A and B monomers, R² is a hydrogen atom or a lower alkyl group, R³ is a lower alkyl group or the residue of a free-radical initiator, n is an integer from 20 to 500 and each R⁴ is a monovalent radical independently selected from the group consisting of



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-CN, and -CO₂R⁶ wherein R⁵ is a hydrogen atom or a lower alkyl group, and R⁶ is a lower alkyl group.

- 15 14. A transdermal drug delivery device according to Claim 1, wherein the macromonomer is selected from the group consisting of polymethylmethacrylate macromonomer, styrene/acrylonitrile macromonomer, and polystyrene macromonomer.
- 20 15. A transdermal drug delivery device according to Claim 1, wherein the softener is present in an amount in excess of 20% and less than about 60% by weight based on the total weight of the matrix.
- 16. A transdermal drug delivery device according to Claim 1, wherein the softener is selected from the group consisting of C₈-C₂₂ fatty acids, C₈-C₂₂ fatty alcohols, lower alkyl esters of C₈-C₂₂ fatty acids, monoglycerides of C₈-C₂₂ fatty acids, di(lower)alkyl esters of C₆-C₈ diacids, tetrahydrofurfuryl alcohol polyethylene glycol ether, polyethylene glycol, propylene glycol, ethoxyethoxy ethanol, diethylene glycol monomethyl ether, N,N-dimethyl dodecylamine-N-oxide, 2-(2-ethoxyethoxy)ethanol, and combinations of the foregoing.

17. A transdermal drug delivery device according to Claim 1, wherein the softener is selected from the group consisting of dimethyl sulfoxide, glycerol, ethanol, ethyl acetate, acetoacetic ester, N-methyl pyrrolidone, isopropyl alcohol, alkylaryl ethers of polyethylene oxide, polyethylene oxide monomethyl ethers, and polyethylene oxide dimethyl ethers.

- 18. A transdermal drug delivery device according to Claim 1, wherein the softener is selected from the group consisting of nicotine, nitroglycerine, chlorpheniramine, nicotinic acid benzyl ester, orphenadrine, scopolamine, and valproic acid.
 - 19. A pressure sensitive skin adhesive comprising:
 - (1) a copolymer comprising

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- (a) one or more A monomers selected from the group

 consisting of alkyl acrylates containing 4 to 10 carbon atoms in the alkyl group and

 alkyl methacrylates containing 4 to 10 carbon atoms in the alkyl group; and
 - (b) optionally one or more ethylenically unsaturated B monomers copolymerizable with the A monomer, and
- (c) a macromonomer copolymerizable with the A and B monomers defined above and having a molecular weight in the range 500-500,000; and
 - (2) a softener dissolved in the copolymer,

wherein the structure and amount of the comonomers in the copolymer, the inherent viscosity of the copolymer, and the amount and structure of the softener are such as to provide the pressure sensitive skin adhesive with a compliance value in the range 2×10^{-6} cm²/dyne to about 4×10^{-3} cm²/dyne.

20. A pressure sensitive skin adhesive according to Claim 19, wherein the B monomer or monomers comprise a functional group selected from the group consisting of carboxylic acid, carboxylic acid ester, hydroxy, sulfonamide, urea, carbamate, carboxamide, amine, oxy, oxo, and cyano.